



# STIC Search Report

## Biotech-Chem Library

STIC Database Tracking Number: 118105

TO: Minh-Tam Davis  
Location: REM/3A24/3C18  
Art Unit: 1642  
Wednesday, March 31, 2004  
Case Serial Number: 10/089600

From: Paul Schulwitz  
Location: Biotech-Chem Library  
REM-1A65  
Phone: (571)272-2527

[paul.schulwitz@uspto.gov](mailto:paul.schulwitz@uspto.gov)

### Search Notes

Examiner Davis,

See attached results.

If you have any questions about this search feel free to contact me at any time.

Thank you for using STIC search services!

Paul Schulwitz  
Technical Information Specialist  
STIC Biotech/Chem Library  
(571)272-2527

*party date 09/99*

STIC-Biotech/ChemLib

118105

From: Chan, Christina  
Sent: Monday, March 29, 2004 3:05 PM  
To: Davis, Minh-Tam; STIC-Biotech/ChemLib  
Subject: RE: Rush search request for 10/089600

Please rush. Thanks Chris

Chris Chan

TC 1600 New Hire Training Coordinator and SPE 1644  
(571)-272-0841  
Remsen, 3E89

-----Original Message-----

From: Davis, Minh-Tam  
Sent: Monday, March 29, 2004 11:49 AM  
To: Chan, Christina  
Subject: Rush search request for 10/089600

Please search in commercial database, issued patent files, PGPUB and interference:  
SEQ ID NO:2.  
Thank you.

MINH TAM DAVIS  
ART UNIT 1642  
ROOM 3A24, MB 3C18  
272-0830

RECEIVED  
MAR 29 2004  
TECH/STIC

Searcher: \_\_\_\_\_  
Phone: \_\_\_\_\_  
Location: \_\_\_\_\_  
Date Picked Up: \_\_\_\_\_  
Date Completed: 3/31  
Searcher Prep/Review: \_\_\_\_\_  
Clerical: \_\_\_\_\_  
Online time: \_\_\_\_\_

TYPE OF SEARCH:  
NA Sequences: \_\_\_\_\_  
AA Sequences: \_\_\_\_\_  
Structures: \_\_\_\_\_  
Bibliographic: \_\_\_\_\_  
Litigation: \_\_\_\_\_  
Full text: \_\_\_\_\_  
Patent Family: \_\_\_\_\_  
Other: \_\_\_\_\_

VENDOR/COST (where applic.)  
STN: \_\_\_\_\_  
DIALOG: \_\_\_\_\_  
Questel/Orbit: \_\_\_\_\_  
DRLink: \_\_\_\_\_  
Lexis/Nexis: \_\_\_\_\_  
Sequence Sys.: \_\_\_\_\_  
WWW/Internet: \_\_\_\_\_  
Other (specify): \_\_\_\_\_





PR 30-SEP-1998; 98US-0102487P.  
 PR 30-SEP-1998; 98US-0102570P.  
 PR 30-SEP-1998; 98US-0102571P.  
 PR 01-OCT-1998; 98US-0102684P.  
 PR 01-OCT-1998; 98US-0102687P.  
 PR 02-OCT-1998; 98US-0102965P.  
 PR 06-OCT-1998; 98US-0103258P.  
 PR 06-OCT-1998; 98US-0103449P.  
 PR 07-OCT-1998; 98US-0103314P.  
 PR 07-OCT-1998; 98US-0103315P.  
 PR 07-OCT-1998; 98US-0103328P.  
 PR 07-OCT-1998; 98US-0103357P.  
 PR 07-OCT-1998; 98US-0103367P.  
 PR 07-OCT-1998; 98US-0103401P.  
 PR 08-OCT-1998; 98US-0103633P.  
 PR 08-OCT-1998; 98US-0103678P.  
 PR 08-OCT-1998; 98US-0103679P.  
 PR 14-OCT-1998; 98US-0104257P.  
 PR 20-OCT-1998; 98US-0104987P.  
 PR 20-OCT-1998; 98US-0105000P.  
 PR 20-OCT-1998; 98US-0105002P.  
 PR 21-OCT-1998; 98US-0105104P.  
 PR 22-OCT-1998; 98US-0105169P.  
 PR 22-OCT-1998; 98US-0105266P.  
 PR 26-OCT-1998; 98US-0105693P.  
 PR 26-OCT-1998; 98US-0105694P.  
 PR 27-OCT-1998; 98US-0105807P.  
 PR 27-OCT-1998; 98US-0105881P.  
 PR 27-OCT-1998; 98US-0105882P.  
 PR 28-OCT-1998; 98US-0106023P.  
 PR 28-OCT-1998; 98US-0106029P.  
 PR 28-OCT-1998; 98US-0106030P.  
 PR 28-OCT-1998; 98US-0106032P.  
 PR 28-OCT-1998; 98US-0106033P.  
 PR 28-OCT-1998; 98US-0106178P.  
 PR 29-OCT-1998; 98US-0106248P.  
 PR 29-OCT-1998; 98US-0106249P.  
 PR 29-OCT-1998; 98US-0106464P.  
 PR 30-OCT-1998; 98US-0106465P.  
 PR 03-NOV-1998; 98US-0106856P.  
 PR 03-NOV-1998; 98US-0106902P.  
 PR 03-NOV-1998; 98US-0106905P.  
 PR 03-NOV-1998; 98US-0106919P.  
 PR 03-NOV-1998; 98US-0106932P.  
 PR 10-NOV-1998; 98US-0106934P.  
 PR 10-NOV-1998; 98US-0107753P.  
 PR 17-NOV-1998; 98US-0108775P.  
 PR 17-NOV-1998; 98US-0108779P.  
 PR 17-NOV-1998; 98US-0108787P.  
 PR 17-NOV-1998; 98US-0108788P.  
 PR 17-NOV-1998; 98US-0108801P.  
 PR 17-NOV-1998; 98US-0108802P.  
 PR 17-NOV-1998; 98US-0108806P.  
 PR 17-NOV-1998; 98US-0108807P.  
 PR 17-NOV-1998; 98US-0108867P.  
 PR 17-NOV-1998; 98US-0108925P.  
 PR 18-NOV-1998; 98US-0108848P.  
 PR 18-NOV-1998; 98US-0108849P.  
 PR 18-NOV-1998; 98US-0108850P.  
 PR 18-NOV-1998; 98US-0108851P.  
 PR 18-NOV-1998; 98US-0108852P.  
 PR 18-NOV-1998; 98US-0108859P.  
 PR 18-NOV-1998; 98US-0108904P.  
 PA (GETH ) GENENTECH INC.  
 XX Baker K, Goddard A, Gurney AL, Smith V, Watanabe CK, Wood WI,  
 FI WFI; 2000-237871/20.  
 DR N-PSDB; AAA37112.  
 XX

PT New mammalian DNA sequences encoding transmembrane, receptor or secreted  
 PT PRO polypeptides, useful for screening of potential peptide or small  
 PT molecule inhibitors of the relevant receptor/ligand interactions.  
 XX  
 XX Claim 12; Fig 182; 773pp; English.  
 XX  
 CC AAA37022 to AAA37144 encode the new isolated human transmembrane,  
 CC receptor or secreted PRO polypeptides given in AA99340 to AA99462. The  
 CC transmembrane and receptor PRO proteins can be used for screening of  
 CC potential peptide or small molecule inhibitors of the relevant  
 CC receptor/ligand interactions. The polypeptides and nucleotide sequences  
 CC encoding them have various industrial applications, including uses as  
 CC pharmaceutical and diagnostic agents. AAA37145 to AAA3730 represent PCR  
 CC primers and hybridisation probes used in the isolation of the PRO  
 CC polypeptides from the present invention  
 XX  
 SQ Sequence 317 AA;  
 Query Match 100.0%; Score 1745; DB 3; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 6; 6e-153;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 1 MARPPENCDCCHLNBAFSSKIKCKISIKGVFGILALTLTLVFMGSKFMPPEPKK 60  
 DB 1 MARPPENCDCCHLNBAFSSKIKCKISIKGVFGILALTLTLVFMGSKFMPPEPKK 60  
 QY 61 AYDMEHTFYSNGEKKIYMEIDPYRTIEIPRSNGTDELEVHDFKNGYTGIVVGLQKC 120  
 DB 61 AYDMEHTFYSNGEKKIYMEIDPYRTIEIPRSNGTDELEVHDFKNGYTGIVVGLQKC 120  
 QY 121 FIKIQIKIIPFSPPEPIDENEITTTTFPQSVIWPAPKPIENRDLKSKILIEICDN 180  
 DB 121 FIKIQIKIIPFSPPEPIDENEITTTTFPQSVIWPAPKPIENRDLKSKILIEICDN 180  
 QY 121 FIKIQIKIIPFSPPEPIDENEITTTTFPQSVIWPAPKPIENRDLKSKILIEICDN 180  
 DB 121 FIKIQIKIIPFSPPEPIDENEITTTTFPQSVIWPAPKPIENRDLKSKILIEICDN 180  
 QY 181 VTMWINPTLISVLELODFEEEGEDLHPANEXKIGIENQEWVPPQVVEKTRRARQASE 240  
 DB 181 VTMWINPTLISVLELODFEEEGEDLHPANEXKIGIENQEWVPPQVVEKTRRARQASE 240  
 QY 241 EELPINDYENGIEPDPMLDRGYCCICYCRGNRYCRVCCEPLIGYVPPYCYGGRYIC 300  
 DB 241 EELPINDYENGIEPDPMLDRGYCCICYCRGNRYCRVCCEPLIGYVPPYCYGGRYIC 300  
 QY 301 RVIMPENWVARMGLGRV 317  
 DB 301 RVIMPENWVARMGLGRV 317  
 RESULT 3  
 AAB25665  
 ID AAB25665 standard; protein; 317 AA.  
 XX  
 XX AAB25665;  
 AC  
 XX 04-DEC-2000 (first entry)  
 DT  
 DE Human secreted protein sequence encoded by gene 1 SEQ ID NO:54.  
 XX  
 XX Human; secreted protein; immunosuppressive; immunostimulant; nocotropic;  
 KW antiinflammatory; cardiant; vulnerary; antitumor; anticonvulsant;  
 KW antiparkinsonian; neuroprotective; antiviral; antibacterial; cyclostatic;  
 KW antiparasitic; thrombolytic; anticoagulant; antidiabetic; cytotoxic;  
 KW gene therapy; vaccine; chemotaxis-modulator; angiogenesis-modulator;  
 KW cancer; immune system disorder; hyperproliferative disorder; infection;  
 KW cardiovascular disorder; neurological disease; wound healing.  
 XX  
 OS Homo sapiens.  
 XX  
 PN WO200043495-A2.  
 XX  
 XX 27-JUL-2000.  
 PD  
 XX 18-JAN-2000; 2000WO-US000903.  
 PF  
 XX

PR 19-JAN-1999; 99US-0116330P.  
 XX  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 XX  
 PI Rosen CA, Ruben SM, Ebner R, Young PE, Ni J, Moore PA;  
 PI Komatsoulis G, Birse CE;  
 XX  
 XX WPI; 2000-499225/44.  
 DR N-PSDB; AAA87666.  
 DR  
 DR  
 PT New isolated polynucleotide encoding a secreted protein useful for  
 PT preventing, treating or ameliorating a medical condition.  
 XX  
 XX Claim 11, Fig 1A-B; 451pp; English.  
 PS  
 PS The polynucleotide sequences given in AAA87666 to AAA87708 encodes the  
 CC human secreted proteins given in AAB25665 to AAB25755. Human secreted  
 CC proteins have activities based on the tissues and cells the genes are  
 CC expressed in. Examples of activities include: immunosuppressive;  
 CC immunostimulant; antiinflammatory; cardiant; vulnary; antitumor;  
 CC neurotropic; antiviral; anticoagulant; antiparasitic; antidiabetic;  
 CC antibacterial; antiparasitic; thrombolytic; anticancer; chemotaxis-  
 CC polynucleotides can be used in gene therapy and as vaccines. Chemotaxis-  
 CC modulators and angiogenesis- modulators. The human secreted proteins and  
 CC polynucleotides can be used for diagnosing (the susceptibility to) a  
 CC pathological condition by determining the presence or absence of a  
 CC mutation in the polynucleotide or determining the presence or amount of  
 CC expression of the protein. The polynucleotides and proteins can also be  
 CC used in the treatment and diagnosis of cancer, diseases of the immune  
 CC system, hyperproliferative disorders, cardiovascular disorders and  
 CC neurological disease. They can also be used to promote wound healing and  
 CC to fight infection. AAA87657 to AAA87665 and AAB25664 represent sequences  
 CC used in the exemplification of the present invention  
 XX  
 SQ Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 3; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 6.6e-153;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCEDCHILNAFAFKSKIKCKSLKICGLVFGILALTLIVFMGSKHMPVPYKK 60  
 DB 1 MAKNPENCEDCHILNAFAFKSKIKCKSLKICGLVFGILALTLIVFMGSKHMPVPYKK 60  
 QY 61 AYDMEHTFYNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
 DB 61 AYDMEHTFYNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
 QY 121 FIKTQIVIPERSEPEBEIDENEITTTFFEQSVIWPAPKPIENRDLKNSKILEICDN 180  
 DB 121 FIKTQIVIPERSEPEBEIDENEITTTFFEQSVIWPAPKPIENRDLKNSKILEICDN 180  
 QY 181 VTMWYINPTLISVSELODFEEGEDLHPANERKKGIEQNEQWVVPQVYKTRHARQASE 240  
 DB 181 VTMWYINPTLISVSELODFEEGEDLHPANERKKGIEQNEQWVVPQVYKTRHARQASE 240  
 QY 241 EELPINDYENGIEFDPMLEDERGYCCICRRGNRYCARVCEPLLGYYPPYCYOGGRVIC 300  
 DB 241 EELPINDYENGIEFDPMLEDERGYCCICRRGNRYCARVCEPLLGYYPPYCYOGGRVIC 300  
 QY 301 RVIMPCNMVAVAMLGKV 317  
 DB 301 RVIMPCNMVAVAMLGKV 317

RESULT 4  
 AAB66179  
 ID AAB66179 standard; protein; 317 AA.  
 XX  
 AC AAB66179;  
 XX  
 DT 02-APR-2001 (first entry)

XX  
 DE Protein of the invention #91.  
 XX  
 XX Secreted; transmembrane; gene therapy.  
 KW  
 XX  
 XX Unidentified.  
 OS  
 XX  
 XX WO200078961-A1.  
 PN  
 XX  
 XX 28-DEC-2000.  
 PD  
 XX  
 XX 18-FEB-2000; 2000WO-US004342.  
 PF  
 XX  
 XX 23-JUN-1999; 99US-0141037P.  
 PR 20-JUL-1999; 99US-0144758P.  
 PR 26-JUL-1999; 99US-0145698P.  
 PR 01-SEP-1999; 99WO-US020111.  
 PR 29-OCT-1999; 99US-0162506P.  
 PR 30-NOV-1999; 99WO-US028313.  
 PR 02-DEC-1999; 99WO-US030551.  
 PR 16-DEC-1999; 99WO-US030095.  
 PR 05-JAN-2000; 2000WO-US000219.  
 PR 06-JAN-2000; 2000WO-US000376.  
 XX  
 XX (GENTH ) GENENTECH INC.  
 XX  
 XX Baker KP, Botstein D, Desnoyers L, Eaton DL, Ferrara N, Fong S;  
 PI Gao W, Goddard A, Godowski PJ, Grimaldi CJ, Guirney AL, Hillan KJ,  
 PI Pan J, Paoni NF, Roy MA, Smith V, Stewart TA, Tumas D, Watanabe CK,  
 PI Williams PM, Wood WJ;  
 XX  
 DR WPI; 2001-071395/08.  
 DR  
 DR Secreted and transmembrane proteins and nucleic acids designated PRO,  
 PT useful as hybridization probes, in chromosome and gene mapping and gene  
 PT therapy.  
 PT  
 PT  
 PS Claim 1, Fig 182; 787pp; English.  
 PS  
 PS The present invention relates to secreted and transmembrane proteins.  
 CC These proteins and the DNA encoding them may be used as hybridization  
 CC probes, in chromosome and gene mapping and in the generation of anti-  
 CC sense RNA and DNA. They may also be used to generate either  
 CC transgenic animals or knockout animals which are in turn useful for  
 CC development and screening of therapeutically useful reagents. The nucleic  
 CC acids may also be used in gene therapy  
 CC  
 XX  
 SQ Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 4; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 6.6e-153;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCEDCHILNAFAFKSKIKCKSLKICGLVFGILALTLIVFMGSKHMPVPYKK 60  
 DB 1 MAKNPENCEDCHILNAFAFKSKIKCKSLKICGLVFGILALTLIVFMGSKHMPVPYKK 60  
 QY 61 AYDMEHTFYNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
 DB 61 AYDMEHTFYNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
 QY 121 FIKTQIVIPERSEPEBEIDENEITTTFFEQSVIWPAPKPIENRDLKNSKILEICDN 180  
 DB 121 FIKTQIVIPERSEPEBEIDENEITTTFFEQSVIWPAPKPIENRDLKNSKILEICDN 180  
 QY 181 VTMWYINPTLISVSELODFEEGEDLHPANERKKGIEQNEQWVVPQVYKTRHARQASE 240  
 DB 181 VTMWYINPTLISVSELODFEEGEDLHPANERKKGIEQNEQWVVPQVYKTRHARQASE 240  
 QY 241 EELPINDYENGIEFDPMLEDERGYCCICRRGNRYCARVCEPLLGYYPPYCYOGGRVIC 300  
 DB 241 EELPINDYENGIEFDPMLEDERGYCCICRRGNRYCARVCEPLLGYYPPYCYOGGRVIC 300

QY 301 RVIMPCNWWVARMGRV 317  
DB 301 RVIMPCNWWVARMGRV 317

## RESULT 5

AA896985  
ID AA896985 standard; protein; 317 AA.

AC AA896985;

DT 27-JUL-2001 (first entry)

DE Human angiogenesis inhibitor related protein SEQ ID NO: 2.

XX Human; mouse; rat; angiogenesis inhibitor; cartilage; Basedow's disease;  
KW diabetic retinopathy; muscular dystrophy; chronic hepatic deficiency;  
KW stomach cancer; chronic bronchitis; osteoarthritis;  
KW chronic rheumatoid arthritis.

OS Homo sapiens.

PN WO200123557-A1.

PD 05-APR-2001.

PF 29-SEP-2000; 2000WO-JP006804.

PR 29-SEP-1999; 99JP-00275947.

PA (TEIJU) TEIJUIN LTD.

XX Yamana K, Takehashi Y, Wada H, Kasahara Y;

XX WPI, 2001-266155/27.

DR N-PSDB; AAF89155.

PT Human, rat and mouse genes encoding angiogenesis inhibitors and cartilage

XX cell differentiating agents.

XX Claim 1; Page 57-58; 84pp; Japanese.

CC The present invention provides the protein and coding sequences of human,  
CC rat and mouse angiogenesis inhibitors. These control the proliferation  
CC and differentiation of cartilage cells and inhibit angiogenesis. They  
CC have potential uses in the treatment of diabetic retinopathy, muscular  
CC dystrophy, Basedow's disease, chronic hepatic deficiency, stomach cancer,  
CC chronic bronchitis, osteoarthritis and chronic rheumatoid arthritis. The  
CC present sequence is a protein given in the exemplification of the  
CC invention

XX SQ Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 4; Length 317;

Best Local Similarity 100.0%; Pred. No. 6, 6e-153;

Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPPNCECHILNNAFAFKKICKSLKICGLVIGILALITIVFWSKHPMPVPPK 60

DB 1 MAKNPPNCECHILNNAFAFKKICKSLKICGLVIGILALITIVFWSKHPMPVPPK 60

QY 61 AYDMEHTFYNGEKKKIYMEIDPVTREIFRSNGGDELTLEVDFFKNGYGIYVGLQKC 120

DB 61 AYDMEHTFYNGEKKKIYMEIDPVTREIFRSNGGDELTLEVDFFKNGYGIYVGLQKC 120

QY 121 FIKTQIVIPFSEPEEIDENEIITTFEQSIVVPAKPIENRDLNKKILICDN 180

DB 121 FIKTQIVIPFSEPEEIDENEIITTFEQSIVVPAKPIENRDLNKKILICDN 180

QY 181 VTMWVNPILISVSELDPFEEGEDIHPANKEKGIQONQWVVPQVKEKTRHAAQASE 240

DB 181 VTMWVNPILISVSELDPFEEGEDIHPANKEKGIQONQWVVPQVKEKTRHAAQASE 240

QY 241 EELPINDYENGIEFPDMLDERGYCCICRGNRVCRAVCEPLIGYPPYCYQGGRAVIC 300  
DB 241 EELPINDYENGIEFPDMLDERGYCCICRGNRVCRAVCEPLIGYPPYCYQGGRAVIC 300  
QY 301 RVIMPCNWWVARMGRV 317  
DB 301 RVIMPCNWWVARMGRV 317

## RESULT 6

AAU12433  
ID AAU12433 standard; protein; 317 AA.

AC AAU12433;

DT 24-OCT-2001 (first entry)

DE Human PRO1565 polypeptide sequence.

XX Human secretory and transmembrane; PRO; mammalian; cancer; lung; breast;  
KW prostate; cervical; tumour necrosis factor-alpha; TNF-alpha; cartilage;  
KW ear; proliferation; glucose; free fatty acid; skeletal muscle; adipocyte;  
KW A-peptide; factor VIIa; gene therapy.

OS Homo sapiens.

PN WO200140466-A2.

PD 07-JUN-2001.

PF 01-DEC-2000; 2000WO-US032678.

PR 01-DEC-1999; 99WO-US028301.

PR 01-DEC-1999; 99WO-US028634.

PR 02-DEC-1999; 99WO-US028551.

PR 02-DEC-1999; 99WO-US028564.

PR 09-DEC-1999; 99WO-US028565.

PR 16-DEC-1999; 99WO-US030095.

PR 20-DEC-1999; 99WO-US030911.

PR 20-DEC-1999; 99WO-US030999.

PR 30-DEC-1999; 99WO-US031243.

PR 30-DEC-1999; 99WO-US031274.

PR 05-JAN-2000; 2000WO-US000277.

PR 06-JAN-2000; 2000WO-US000376.

PR 06-JAN-2000; 2000WO-US003565.

PR 11-FEB-2000; 2000WO-US004341.

PR 18-FEB-2000; 2000WO-US004342.

PR 22-FEB-2000; 2000WO-US004414.

PR 24-FEB-2000; 2000WO-US004914.

PR 01-MAR-2000; 2000WO-US005601.

PR 02-MAR-2000; 2000WO-US005841.

PR 03-MAR-2000; 2000US-0187202P.

PR 10-MAR-2000; 2000WO-US006319.

PR 15-MAR-2000; 2000WO-US006884.

PR 20-MAR-2000; 2000WO-US007377.

PR 21-MAR-2000; 2000WO-US007532.

PR 30-MAR-2000; 2000WO-US008439.

PR 17-MAY-2000; 2000WO-US013705.

PR 22-MAY-2000; 2000WO-US014042.

PR 30-MAY-2000; 2000WO-US014941.

PR 02-JUN-2000; 2000WO-US015264.

PR 05-JUN-2000; 2000US-0209832P.

PR 28-JUL-2000; 2000WO-US020710.

PR 11-AUG-2000; 2000WO-US022031.

PR 23-AUG-2000; 2000WO-US023522.

PR 24-AUG-2000; 2000WO-US023528.

PR 08-NOV-2000; 2000WO-US030952.

PR 10-NOV-2000; 2000WO-US030873.

PA (GETH) GENENTECH INC.

XX Baker KP, Beresini M, DeForge L, Desnoyers L, Filvaroff E, Gao W,  
PI Gerritsen ME, Goddard A, Godowski PJ, Gurney AL, Sherwood S;  
PI Smith V, Stewart TA, Tumas D, Watanabe CK, Wood WI, Zhang Z;  
XX WPI, 2001:408281/43.  
DR N-PSDB; AAS21505.  
XX Isolated, secretory and transmembrane PRO polypeptide used to detect  
PT other PRO polypeptides, link bioactive molecules to cells expressing PRO  
PT polypeptides, and detect the presence of mammalian tumors e.g. lung,  
PT breast, prostate, cervical.

XX Claim 12, Fig 524; 813pp; English.

XX AAU12172-AAU12446 represent novel human secretory and transmembrane PRO  
CC polypeptides. The PRO polypeptides are useful to detect other PRO  
CC polypeptides, to link bioactive molecules to cells expressing PRO  
CC polypeptides, to modulate biological activities of cells expressing PRO  
CC polypeptides, and to detect the presence of mammalian lung, colon,  
CC breast, prostate, rectal, cervical or liver tumors by comparing PRO  
CC polypeptide expression in a cell sample to that in a control sample. Some  
CC of the 275 sequences are also useful to stimulate the release of tumor  
CC necrosis factor-alpha (TNF-alpha) from human blood, the proliferation or  
CC differentiation of chondrocytes, the proliferation or gene expression in  
CC pericyte cells, the release of proteoglycans from cartilage, the  
CC proliferation of inner ear utricular supporting cells or of T-  
CC lymphocytes, the release of a cytokine from peripheral blood monocytes  
CC (PBMCs), or the proliferation of endothelial cells. Some of the PRO  
CC polypeptides may modulate glucose or free fatty acid uptake by skeletal  
CC muscle cells or by adipocytes; or inhibit binding of A-peptide to factor  
CC VIIA. The PRO polypeptides can be used in assays to identify molecules  
CC involved in binding interactions. The polynucleotides encoding PRO  
CC polypeptides can be used to generate probes, antisense RNA/DNA,  
CC transgenic or knock out animals and can be used in gene therapy

XX Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 4; Length 317;  
Best Local Similarity 100.0%; Pred. No. 6.6e-153; Indels 0; Gaps 0;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKAPPENECEDCHILNAEAFKSKKICSKLTCGLVFGILATLIVLFGSKHFWPEVPKX 60  
DB 1 MAKAPPENECEDCHILNAEAFKSKKICSKLTCGLVFGILATLIVLFGSKHFWPEVPKX 60  
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTREIFRSGNGTDETLVHDFKNGYTGIFYVGLQKC 120  
DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTREIFRSGNGTDETLVHDFKNGYTGIFYVGLQKC 120  
QY 121 FIKTQIKVLPFSEPEEIDEENEITTTFFEQSVIWPAPKEPIENRDFLNSKILIEICDN 180  
DB 121 FIKTQIKVLPFSEPEEIDEENEITTTFFEQSVIWPAPKEPIENRDFLNSKILIEICDN 180  
QY 181 VTMWVINTLLISVSLQDFEEGEDLHPANKEKGIENQEWVVPQVKVETRRARQASE 240  
DB 181 VTMWVINTLLISVSLQDFEEGEDLHPANKEKGIENQEWVVPQVKVETRRARQASE 240  
QY 241 EELPINDYTGIEEDPMLDERGYCCICRGRNRYCRVCEPPLIGYPPYCYOGGAVIC 300  
DB 241 EELPINDYTGIEEDPMLDERGYCCICRGRNRYCRVCEPPLIGYPPYCYOGGAVIC 300  
QY 301 RVIMPQNMVAVARMLGRV 317  
DB 301 RVIMPQNMVAVARMLGRV 317

RESULT 7  
AAB87583  
ID AAB87583 standard; protein; 317 AA.

XX AAB87583;  
XX

DT 15-MAY-2001 (first entry)  
XX Human PRO1565.  
DB Human PRO1565.  
XX Human, PRO protein; mapping.  
XX Homo sapiens.  
OS Homo sapiens.  
XX WO200116318-A2.  
EN 08-MAR-2001.  
PD 24-AUG-2000; 2000WO-US023328.  
PF 01-SEP-1999; 99WO-US020111.  
PR 15-SEP-1999; 99WO-US021030.  
PR 07-DEC-1999; 99US-0169495P.  
PR 09-DEC-1999; 99US-0170262P.  
PR 11-JUN-2000; 2000US-0175481P.  
PR 18-FEB-2000; 2000WO-US004341.  
PR 18-FEB-2000; 2000WO-US004342.  
PR 22-FEB-2000; 2000WO-US004414.  
PR 01-MAR-2000; 2000WO-US005601.  
PR 23-MAR-2000; 2000US-0187202P.  
PR 21-MAR-2000; 2000US-0191007P.  
PR 30-MAR-2000; 2000WO-US008439.  
PR 25-APR-2000; 2000US-0199397P.  
PR 22-MAY-2000; 2000WO-US014042.  
PR 05-JUN-2000; 2000US-0209832P.

XX (GENT) GENENTECH INC.

PI Eaton DL, Filvaroff E, Gerritsen ME, Goddard A, Godowski PJ;  
PI Grimaldi CJ, Gurney AL, Watanabe CK, Wood WI;  
DR N-PSDB; AAF92115.  
XX WPI, 2001-183260/18.

XX Eighty four nucleic acids encoding PRO polypeptides, useful in molecular  
PT biology, including use as hybridization probes, and in chromosome and  
PT gene mapping.

XX Claim 12; Fig 116; 278pp; English.

XX The present sequence is a human PRO polypeptide (secreted and  
CC transmembrane). The PRO protein, and PRO agonists, PRO antagonists or  
CC anti-PRO antibodies are useful for preparation of a medicament useful in  
CC the treatment of a condition which is responsive to the PRO protein,  
CC agonists, antagonists or anti-PRO antibodies. The PRO protein may also be  
CC employed as molecular weight markers for protein electrophoresis. The PRO  
CC coding sequence has applications in molecular biology, including use as  
CC hybridization probes, and in chromosome and gene mapping

XX Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 4; Length 317;  
Best Local Similarity 100.0%; Pred. No. 6.6e-153; Indels 0; Gaps 0;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKAPPENECEDCHILNAEAFKSKKICSKLTCGLVFGILATLIVLFGSKHFWPEVPKX 60  
DB 1 MAKAPPENECEDCHILNAEAFKSKKICSKLTCGLVFGILATLIVLFGSKHFWPEVPKX 60  
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTREIFRSGNGTDETLVHDFKNGYTGIFYVGLQKC 120  
DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTREIFRSGNGTDETLVHDFKNGYTGIFYVGLQKC 120  
QY 121 FIKTQIKVLPFSEPEEIDEENEITTTFFEQSVIWPAPKEPIENRDFLNSKILIEICDN 180  
DB 121 FIKTQIKVLPFSEPEEIDEENEITTTFFEQSVIWPAPKEPIENRDFLNSKILIEICDN 180  
QY 181 VTMWVINTLLISVSLQDFEEGEDLHPANKEKGIENQEWVVPQVKVETRRARQASE 240  
DB 181 VTMWVINTLLISVSLQDFEEGEDLHPANKEKGIENQEWVVPQVKVETRRARQASE 240



Db 181 VTMWVNPPLISVSELDFFESEGEDLHPANKEKGIQNEQWVPQVKEKTRHARQASE 240

QY 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRCRRVCEPPLGGYPPYCYGGGRVYC 300

Db 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRCRRVCEPPLGGYPPYCYGGGRVYC 300

QY 301 RVIMPCNMVWVARMIGRV 317

Db 301 RVIMPCNMVWVARMIGRV 317

RESULT 8

AAB9936 standard; protein; 317 AA.

AAB9936; 27-SEP-2001 (first entry)

XX Human new chondromodulin-I (NCHM) protein SEQ ID NO:1.

XX Human; new chondromodulin-I; NCHM; CHM-I; bone; cartilage; joint; cancer;

XX malignant tumour; pathologic angiogenesis; organ insufficiency;

XX digestive tract failure; exocrine secretion failure; osteopathic;

XX cytostatic; cardiovascular; gastrointestinal; abnormal vascularisation;

XX foetal growth disorder; organ failure; digestive disorder;

XX secretory disorder.

XX Homo sapiens.

XX WO200148203-A1.

XX 05-JUL-2001.

XX 26-DEC-2000; 2000WO-JP009236.

XX 27-DEC-1999; 99JP-00369946.

XX 17-FEB-2000; 2000JP-00045124.

XX (TAKE) TAKEDA CHEM IND LTD.

XX Itoh Y, Nishi K;

XX WPI: 2001-425668/45.

XX N-PSDB: AAH44410, AAH44417.

XX New proteins for treating bone, cartilage and joint diseases, cancer,

XX abnormal vascularisation, fetal growth disorders, organ failure,

XX digestive disorders and secretory disorders.

XX Claim 1; Page 69-71; 83pp; Japanese.

XX The present sequence represents the human new chondromodulin-I (NCHM)

XX protein. NCHM has osteopathic, cytostatic, cardiovascular and

XX gastrointestinal activities. The NCHM protein can be used for the

XX treatment and prevention of bone, cartilage and joint diseases, cancer,

XX abnormal vascularisation, foetal growth disorders, organ failure,

XX digestive disorders and secretory disorders

XX Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 4; Length 317;

Best Local Similarity 100.0%; Pred. No. 6.6e-153; Indels 0; Gaps 0;

Matches 317; Conservative 0; Mismatches 0;

QY 1 MAKXPENCECHILNNAFAFKSKICKSICIGLVFGILALTLLVLFWGSKHFWPEVPKX 60

Db 1 MAKXPENCECHILNNAFAFKSKICKSICIGLVFGILALTLLVLFWGSKHFWPEVPKX 60

QY 61 AYDMHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKXGYTGIVFVGLQKC 120

Db 61 AYDMHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKXGYTGIVFVGLQKC 120

QY 121 FIKTQIKVIFPFSRPEREIDENERITTTFFEQSVIWPAPKEPIENRDLKSKLLEICDN 180

Db 121 FIKTQIKVIFPFSRPEREIDENERITTTFFEQSVIWPAPKEPIENRDLKSKLLEICDN 180

QY 181 VTMWVNPPLISVSELDFFESEGEDLHPANKEKGIQNEQWVPQVKEKTRHARQASE 240

Db 181 VTMWVNPPLISVSELDFFESEGEDLHPANKEKGIQNEQWVPQVKEKTRHARQASE 240

QY 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRCRRVCEPPLGGYPPYCYGGGRVYC 300

Db 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRCRRVCEPPLGGYPPYCYGGGRVYC 300

QY 301 RVIMPCNMVWVARMIGRV 317

Db 301 RVIMPCNMVWVARMIGRV 317

RESULT 9

AAB82755 standard; protein; 317 AA.

XX AAB82755; 29-OCT-2001 (first entry)

XX Human chondromodulin-I related peptide.

XX Chondromodulin-I related peptide; ChMIRp; human; tendon; muscle;

XX cartilage; bone; tendinitis; cachexia; muscular dystrophy; inflammation;

XX allergy; wound healing; arthritis; infertility; vulnerability; diagnosis;

XX antiinflammatory; antiallergic; antiarthritic; osteogenic; diagnosis;

XX therapy.

XX Homo sapiens.

XX Location/Qualifiers

XX Key 276

XX Misc-difference /note= "Cys-276 may be replaced by Ser or Ala in a ChMIRp

XX polypeptide of Claim 17"

XX Misc-difference 280

XX /note= "Cys-280 may be replaced by Ser or Ala in a ChMIRp

XX polypeptide of Claim 18"

XX Misc-difference 281

XX /note= "Glu-281 may be replaced by Asp in a ChMIRp

XX polypeptide of Claim 19"

XX Misc-difference 285

XX /note= "Gly-285 may be replaced by Pro or Ala in a ChMIRp

XX polypeptide of Claim 20"

XX Misc-difference 297

XX /note= "Arg-297 may be replaced by Lys, Gln or Asn in a

XX ChMIRp polypeptide of Claim 21"

XX Misc-difference 300

XX /note= "Cys-300 may be replaced by Ser or Ala in a ChMIRp

XX polypeptide of Claim 22"

XX Misc-difference 306

XX /note= "Cys-306 may be replaced by Ser or Ala in a ChMIRp

XX polypeptide of Claim 23"

XX Misc-difference 310

XX /note= "Val-310 may be replaced by Ile, Met, Leu, Phe,

XX Ala or Nle in a ChMIRp polypeptide of Claim 24"

XX MO20015344-A2.

XX 26-UTL-2001.

XX 18-JAN-2001; 2001WO-US001700.

XX 19-JAN-2000; 2000US-0176898P.

XX 28-NOV-2000; 2000US-00724310.

XX (AMGE-) AMGEN INC.

XX Nguyen H, Jung J, Clarkin K;

XX WPI: 2001-536401/59.  
DR N-PSDB; AAH26446; AAH26448.  
XX New chondromodulin-I like polypeptides and polynucleotides useful for  
PT treating, preventing or ameliorating diseases resulting from abnormal  
PT levels of ChnMtrp, or for inducing cartilage formation and bone growth.  
XX Claim 14; Fig 2; 189pp; English.  
PS  
XX  
CC The present sequence is that of human chondromodulin-I related peptide  
CC (ChnMtrp), as predicted from the open reading frame of an isolated cDNA  
CC clone (see AAH26446). The protein is structurally related to  
CC chondromodulin-I. It is expressed in the tendons, skeletal muscle,  
CC thymus, ovary, cerebral cortex, intestine M cells and in cells adjacent  
CC to the hair follicle. The invention provides ChnMtrp polypeptides  
CC (including those having the present sequence, or at least 1 conservative  
CC amino acid substitution, an insertion or deletion, or C- and/or N-  
CC terminal truncation, while maintaining activity) and polynucleotides, as  
CC well as vectors, host cells, antibodies and recombinant methods for  
CC producing the polypeptide. It also discloses therapeutic and diagnostic  
CC uses for these and related products. ChnMtrp may play a role in cartilage  
CC development and blood vessel formation, in the development of tendons and  
CC muscle and in attachment of muscle to bone, in immune function, and as a  
CC growth factor in the growth and development of various tissues. It may be  
CC useful for the diagnosis and/or treatment of tendon diseases (e.g.  
CC tendinitis and tendon tear), skeletal muscle diseases (e.g. cachexia and  
CC muscular dystrophy), immune system dysfunction diseases (e.g.  
CC inflammation, allergy, poor wound healing and arthritis) and infertility  
CC diseases  
CC  
SQ Sequence 317 AA;  
Query Match 100.0%; Score 1745; DB 4; Length 317;  
Best Local Similarity 100.0%; Pred. No. 6,6e-153;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 1 MAKNPENCECHINAAEFKSKTCKSLKTCGLVGLIALTLVLFMGSHFPEVPK 60  
DB 1 MAKNPENCECHINAAEFKSKTCKSLKTCGLVGLIALTLVLFMGSHFPEVPK 60  
QY 61 AYDMHTFVSNKSKKIMWEIDPVTRTEIPRSGNGTDETLKHPKNGYTIYVGLQKC 120  
DB 61 AYDMHTFVSNKSKKIMWEIDPVTRTEIPRSGNGTDETLKHPKNGYTIYVGLQKC 120  
QY 121 PIKTOIKVLPFSEPEEIDEENEITTTFFEQSVIWPAAKPIENRDFLKNKSLIETCDN 180  
DB 121 PIKTOIKVLPFSEPEEIDEENEITTTFFEQSVIWPAAKPIENRDFLKNKSLIETCDN 180  
QY 181 VTMWAINPTLISVSELODFEESGDLHPANKEKGIENQWVVPQVVEKTRARQSE 240  
DB 181 VTMWAINPTLISVSELODFEESGDLHPANKEKGIENQWVVPQVVEKTRARQSE 240  
QY 241 EELPIINDYENGIEFPDMLDERGVCICRCRNGYCRVCEPCELLGYPPYCYOGGRVTC 300  
DB 241 EELPIINDYENGIEFPDMLDERGVCICRCRNGYCRVCEPCELLGYPPYCYOGGRVTC 300  
QY 301 RVIMPONWVARMIGRV 317  
DB 301 RVIMPONWVARMIGRV 317  
RESULT 10  
ABG95908  
ID ABG95908 standard; protein; 317 AA.  
AC ABG95908;  
XX  
XX  
DT 10-DEC-2002 (fixed entry)  
XX  
XX Human secreted/transmembrane protein PRO1565.  
DE  
XX Human; secreted protein; transmembrane protein; antirheumatic;  
KM

KW antiarthritis; osteopathic; sports-related joint problem;  
KM articular cartilage defect; osteoarthritis; rheumatoid arthritis.  
XX  
XX Homo sapiens.  
OS  
XX US2002119130-A1.  
XX  
PD 29-AUG-2002.  
XX  
XX 06-DEC-2001; 2001US-00006867.  
XX  
XX 29-OCT-1997; 97US-0063435F.  
XX 29-OCT-1997; 97US-0064215F.  
XX 22-APR-1998; 98US-0082797F.  
XX 29-APR-1998; 98US-0083495F.  
XX 15-MAY-1998; 98US-0085579F.  
XX 02-JUN-1998; 98US-008759F.  
XX 04-JUN-1998; 98US-0088021F.  
XX 04-JUN-1998; 98US-0088029F.  
XX 04-JUN-1998; 98US-0088030F.  
XX 10-JUN-1998; 98US-0088734F.  
XX 10-JUN-1998; 98US-0088740F.  
XX 10-JUN-1998; 98US-0088811F.  
XX 10-JUN-1998; 98US-0088824F.  
XX 10-JUN-1998; 98US-0088825F.  
XX 11-JUN-1998; 98US-0088835F.  
XX 12-JUN-1998; 98US-0089105F.  
XX 16-JUN-1998; 98US-0089514F.  
XX 17-JUN-1998; 98US-0089531F.  
XX 19-JUN-1998; 98US-0089952F.  
XX 22-JUN-1998; 98US-0090246F.  
XX 24-JUN-1998; 98US-0090444F.  
XX 25-JUN-1998; 98US-0090688F.  
XX 25-JUN-1998; 98US-0090696F.  
XX 26-JUN-1998; 98US-0090862F.  
XX 02-JUL-1998; 98US-0091628F.  
XX 10-AUG-1998; 98US-0096012F.  
XX 17-AUG-1998; 98US-0096757F.  
XX 18-AUG-1998; 98US-0096849F.  
XX 18-AUG-1998; 98US-0096959F.  
XX 26-AUG-1998; 98US-0097954F.  
XX 26-AUG-1998; 98US-0097971F.  
XX 26-AUG-1998; 98US-0097979F.  
XX 01-SEP-1998; 98US-0098749F.  
XX 10-SEP-1998; 98US-0099741F.  
XX 10-SEP-1998; 98US-0099763F.  
XX 10-SEP-1998; 98US-0099792F.  
XX 10-SEP-1998; 98US-0099812F.  
XX 10-SEP-1998; 98US-0099815F.  
XX 16-SEP-1998; 98US-0100627F.  
XX 16-SEP-1998; 98US-0100662F.  
XX 16-SEP-1998; 98US-01019330.  
XX 17-SEP-1998; 98US-0100683F.  
XX 17-SEP-1998; 98US-0100684F.  
XX 17-SEP-1998; 98US-0100930F.  
XX 22-SEP-1998; 98US-0101279F.  
XX 23-SEP-1998; 98US-0101475F.  
XX 24-SEP-1998; 98US-0101738F.  
XX 24-SEP-1998; 98US-0101743F.  
XX 24-SEP-1998; 98US-0101916F.  
XX 24-SEP-1998; 98US-0102570F.  
XX 30-SEP-1998; 98US-0103449F.  
XX 06-OCT-1998; 98US-0103449F.  
XX 08-MAR-1999; 99WO-US005028.  
XX 14-MAY-1999; 99WO-US010733.  
XX 02-JUN-1999; 99WO-US012252.  
XX 01-SEP-1999; 99WO-US020111.  
XX 15-SEP-1999; 99WO-US021090.  
XX 15-SEP-1999; 99WO-US021194.  
XX 22-DEC-1999; 99WO-US030720.  
XX 18-FEB-2000; 2000WO-US004341.  
XX 18-FEB-2000; 2000WO-US004342.  
XX 22-FEB-2000; 2000WO-US004414.  
XX 01-MAR-2000; 2000WO-US005601.  
PR

30-MAR-2000; 2000WO-US008439.  
 22-MAY-2000; 2000WO-US014042.  
 PR 02-JUN-2000; 2000WO-US015264.  
 PR 23-AUG-2000; 2000WO-US023522.  
 PR 24-AUG-2000; 2000WO-US023328.  
 PR 10-NOV-2000; 2000WO-US030873.  
 PR 01-DEC-2000; 2000WO-US032378.  
 PR 20-DEC-2000; 2000WO-US034956.  
 PR 28-FEB-2001; 2001WO-US006520.  
 PR 01-MAR-2001; 2001WO-US006666.  
 PR 30-MAY-2001; 2001WO-US017443.  
 PR 01-JUN-2001; 2001WO-US017800.  
 PR 20-JUN-2001; 2001WO-US019692.  
 PR 29-JUN-2001; 2001WO-US021066.  
 PR 09-JUL-2001; 2001WO-US021735.  
 XX (GETH ) GENENTECH INC.  
 XX Eaton DL, Filvaroff E, Gerritsen ME, Goddard A, Godowski PJ;  
 PI Grimaldi JC, Gurney AL, Matanabe CK, Wood MI;  
 DR MPI; 2002-71348/79.  
 XX N-PSDB; ABS74435.  
 PT New isolated secreted and transmembrane PRO polypeptide useful for  
 PT modulating biological activity of a cell, or for treating sports-related  
 PT joint problems, osteoarthritis or rheumatoid arthritis.  
 XX Claim 20; Fig 116; 399pp; English.  
 XX The invention relates to an isolated secreted and transmembrane PRO  
 CC polypeptide having 80 % sequence identity to a sequence appearing as  
 CC ABG95851-ABG95934 or their associated signal peptide, or a sequence of an  
 CC extracellular domain of the proteins with their associated signal peptide  
 CC or lacking its associated signal peptide. Also included are the nucleic  
 CC acids encoding the proteins, vectors, host cells, fusion proteins and  
 CC antibodies which specifically bind to the proteins. The proteins are  
 CC useful for detecting a polypeptide designated as A, B, C or D in a sample  
 CC suspected of containing an A, B, C or D polypeptide, by contacting the  
 CC sample with a polypeptide designated as E, F, G, H or I (or vice versa)  
 CC and determining the formation of a A/E, B/F, B/G, C/H or D/I polypeptide  
 CC conjugate in the sample, where the formation of the conjugate is  
 CC indicative of the presence of an A, B, C or D polypeptide in the sample,  
 CC where A is a PRO10272 polypeptide, B is a PRO20110 polypeptide, C is a  
 CC PRO10096 polypeptide, D is a PRO19760 polypeptide, E is a PRO5801  
 CC polypeptide, F is a PRO1 polypeptide, G is a PRO20040 polypeptide, H is a  
 CC PRO20233 polypeptide and I is a PRO1899 polypeptide. The sample comprises  
 CC a cell suspected of expressing the A, B, C or D polypeptide. The E, F, G,  
 CC H or I polypeptide is labeled with a detectable label or is attached to a  
 CC solid support. The proteins are useful for linking a bioactive molecule  
 CC to a cell expressing a polypeptide designated as A, B, C or D or E, F, G,  
 CC H or I. The bioactive molecule is a toxin, a radiolabel or an antibody.  
 CC The bioactive molecule causes death of the cell. A, B, C, D, E, F, G, H,  
 CC or I, or antibodies against them are useful for modulating a biological  
 CC activity of a cell expressing a polypeptide designated as A, B, C or D or  
 CC E, F, G, H, or I. The cell is killed. The proteins are useful for  
 CC identifying agonists or antagonists, for the preparation of a medicament  
 CC useful in the treatment of a condition which is responsive to the  
 CC proteins, as molecular weight markers for protein electrophoresis  
 CC purposes, as therapeutic agents for treating sports-related joint  
 CC problems, articular cartilage defects, osteoarthritis or rheumatoid  
 CC arthritis. Nucleic acids encoding the proteins are useful as  
 CC hybridisation probes, in chromosome and gene mapping, in the generation  
 CC of anti-sense RNA and DNA, for the preparation of the proteins, to  
 CC generate transgenic or knockout animals which are useful in the  
 CC development and screening of therapeutic useful reagents, for chromosome  
 CC identification, and in gene therapy. The antibody is useful as a  
 CC therapeutic agent, in a diagnostic assay and for affinity purification of  
 CC the protein from recombinant cell culture natural sources. The present  
 CC sequence represents a novel secreted or transmembrane protein of the  
 CC invention  
 XX Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 5; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 6, 6e-153; Indels 0; Gaps 0;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 1 MAKPPEPCEDCHILNAEAFKSKKIKGVLFGIILALIVLFWGSKHFWPEVPPK 60  
 Db 1 MAKPPEPCEDCHILNAEAFKSKKIKGVLFGIILALIVLFWGSKHFWPEVPPK 60  
 QY 61 AYMEHTFYNSGKKKIKYIMIDPVTTELFRSNGNDETELVDFKNGYGIYFVGLQKC 120  
 Db 61 AYMEHTFYNSGKKKIKYIMIDPVTTELFRSNGNDETELVDFKNGYGIYFVGLQKC 120  
 QY 121 FIKQIVIPESFPESEIDENEITTTTFEQSVIWPAPKPIENRDFLNKSKILBICDN 180  
 Db 121 FIKQIVIPESFPESEIDENEITTTTFEQSVIWPAPKPIENRDFLNKSKILBICDN 180  
 QY 181 VTWVWNPFLISVSELDPEEGEDLHFPANERKGIENQWVWPQVYKVKETHARQASE 240  
 Db 181 VTWVWNPFLISVSELDPEEGEDLHFPANERKGIENQWVWPQVYKVKETHARQASE 240  
 QY 241 EELPINDYENGJEPFMDLDERGVCICRCRGNRCRVCEPPLGGYPYCYOGGRVIC 300  
 Db 241 EELPINDYENGJEPFMDLDERGVCICRCRGNRCRVCEPPLGGYPYCYOGGRVIC 300  
 QY 301 RVIMPCKMWWYARMIGRY 317  
 Db 301 RVIMPCKMWWYARMIGRY 317  
 RESULT 11  
 ID ABO17877 standard; protein; 317 AA.  
 AC ABO17877;  
 XX 26-AUG-2003 (first entry)  
 XX DE Novel human secreted and transmembrane protein PRO1565.  
 XX Human; secreted and transmembrane protein; PRO; antiinflammatory;  
 KW antiarteriosclerotic; cardiant; anti-infectility; anti-HIV; cytostatic;  
 KW antidiabetic; gene therapy; tumour necrosis factor (TNF)-alpha release;  
 KW TNF-alpha release; cell proliferation; cell differentiation;  
 KW gene expression modulator; proteoglycan release; cytokine release;  
 KW tumour; inflammatory disease; organ failure; atherosclerosis;  
 KW cardiac injury; infectility; birth defect; premature aging; AIDS;  
 KW acquired immunodeficiency syndrome; cancer; diabetic complication;  
 KW chromosome mapping; gene mapping; pharmaceutical; diagnostic; biosensor;  
 KW bioreactor; tissue typing.  
 OS Homo sapiens.  
 XX US2003032156-A1.  
 XX 13-FEB-2003.  
 XX 06-MAY-2002; 2002US-00140474.  
 XX 31-MAR-1997; 97WO-US005230.  
 PR 12-JUN-1998; 98WO-US012456.  
 PR 14-JUL-1998; 98WO-US014552.  
 PR 28-AUG-1998; 98WO-US017868.  
 PR 10-SEP-1998; 98WO-US018824.  
 PR 14-SEP-1998; 98WO-US019093.  
 PR 14-SEP-1998; 98WO-US019093.  
 PR 14-SEP-1998; 98WO-US019177.  
 PR 14-SEP-1998; 98WO-US019330.  
 PR 16-SEP-1998; 98WO-US019437.  
 PR 17-SEP-1998; 98WO-US021141.  
 PR 07-OCT-1998; 98WO-US022991.  
 PR 29-OCT-1998; 98WO-US022992.  
 PR 20-NOV-1998; 98WO-US024855.

PR	05-JAN-1998	99MO-US0250108
PR	05-JAN-1999	99MO-US0250106
PR	10-MAR-1999	99MO-US0050528
PR	10-MAR-1999	99MO-US0050519
PR	20-MAR-1999	99MO-US0086815
PR	14-MAY-1999	99MO-US0107033
PR	02-JUN-1999	99MO-US012252
PR	01-SEP-1999	99MO-US0201111
PR	08-SEP-1999	99MO-US020594
PR	13-SEP-1999	99MO-US020944
PR	15-SEP-1999	99MO-US021099
PR	15-SEP-1999	99MO-US021547
PR	05-OCT-1999	99MO-US023088
PR	29-NOV-1999	99MO-US028214
PR	30-NOV-1999	99MO-US028313
PR	30-NOV-1999	99MO-US028409
PR	01-DEC-1999	99MO-US0283001
PR	01-DEC-1999	99MO-US028554
PR	02-DEC-1999	99MO-US028564
PR	02-DEC-1999	99MO-US028565
PR	16-DEC-1999	99MO-US030095
PR	20-DEC-1999	99MO-US030911
PR	22-DEC-1999	99MO-US030999
PR	22-DEC-1999	99MO-US030720
PR	30-DEC-1999	99MO-US031243
PR	30-DEC-1999	99MO-US031274
PR	05-JAN-2000	2000MO-US000219
PR	06-JAN-2000	2000MO-US000277
PR	06-JAN-2000	2000MO-US000376
PR	11-FEB-2000	2000MO-US003565
PR	18-FEB-2000	2000MO-US003441
PR	18-FEB-2000	2000MO-US003442
PR	22-FEB-2000	2000MO-US004143
PR	24-FEB-2000	2000MO-US004914
PR	01-MAR-2000	2000MO-US005504
PR	01-MAR-2000	2000MO-US005601
PR	02-MAR-2000	2000MO-US005746
PR	02-MAR-2000	2000MO-US005841
PR	15-MAR-2000	2000MO-US006319
PR	15-MAR-2000	2000MO-US006884
PR	21-MAR-2000	2000MO-US007337
PR	17-MAY-2000	2000MO-US013705
PR	22-MAY-2000	2000MO-US014042
PR	30-MAY-2000	2000MO-US019411
PR	02-JUN-2000	2000MO-US019411
PR	28-JUL-2000	2000MO-US020710
PR	11-AUG-2000	2000MO-US022031
PR	23-AUG-2000	2000MO-US023522
PR	24-AUG-2000	2000MO-US023328
PR	08-NOV-2000	2000MO-US030952
PR	10-NOV-2000	2000MO-US030873
PR	10-DEC-2000	2000MO-US032678
PR	20-DEC-2000	2000MO-US047259
PR	28-FEB-2001	2000MO-US049556
PR	28-FEB-2001	2001MO-US076498
PR	01-MAR-2001	2001MO-US006650
PR	09-MAR-2001	2001MO-US007052
PR	14-MAR-2001	2001US-US0086869
PR	22-MAR-2001	2001US-US0816744
PR	05-APR-2001	2001US-US0828166
PR	10-MAY-2001	2001US-US0854088
PR	18-MAY-2001	2001US-US0854880
PR	25-MAY-2001	2001US-US0860216
PR	25-MAY-2001	2001US-US0866034
PR	01-JUN-2001	2001US-US0872035
PR	01-JUN-2001	2001MO-US0178052
PR	05-JUN-2001	2001US-US0178053

[illegible]

OY 241 BELPINDYENGIEFDPMIDERGVCCTICRGGRNRCRVCEPLIGYPPYCYGGGRVIC 300  
DB 241 BELPINDYENGIEFDPMIDERGVCCTICRGGRNRCRVCEPLIGYPPYCYGGGRVIC 300  
OY 301 RVINECNMMVARMIGRV 317  
DB 301 RVINECNMMVARMIGRV 317

RESULT 12  
ABU81131  
ID ABU81131 standard; protein; 317 AA.  
XX  
AC ABU81131;  
XX  
DT 23-JUN-2003 (first entry)  
XX  
DE Human PRO polypeptide #262.  
XX  
KW Human; PRO polypeptide; secreted and transmembrane protein;  
KW anti-PRO antibody; diagnostic assay; gene expression; diabetes;  
KW bone disorder; carriage disorder; rheumatoid arthritis; obesity;  
KW sports injury; osteoarthritis; hyper-insulinemia; hypo-insulinemia;  
KW hearing loss; coagulation disorder; stroke; heart attack; cardiac;  
KW antidiabetic; anorectic; vulnerable; antiarthritic; osteopathic;  
KW antihemetic; auditory; cerebroprotective; angiogenic.  
XX  
OS Homo sapiens.  
XX  
PN US2003004311-A1.  
XX  
PD 02-JAN-2003.  
XX  
PF 19-DEC-2001; 2001US-00028072.  
XX  
PR 18-JUN-1997; 97US-0049911P.  
PR 26-AUG-1997; 97US-0056974P.  
PR 17-SEP-1997; 97US-0059113P.  
PR 17-SEP-1997; 97US-0059115P.  
PR 17-SEP-1997; 97US-0059117P.  
PR 17-SEP-1997; 97US-0059122P.  
PR 17-SEP-1997; 97US-0059184P.  
PR 18-SEP-1997; 97US-0059263P.  
PR 19-SEP-1997; 97US-0059352P.  
PR 19-SEP-1997; 97US-0059388P.  
PR 24-SEP-1997; 97US-0059836P.  
PR 17-OCT-1997; 97US-0062285P.  
PR 17-OCT-1997; 97US-0062287P.  
PR 17-OCT-1997; 97US-0063755P.  
PR 17-OCT-1997; 97US-0062814P.  
PR 24-OCT-1997; 97US-0062816P.  
PR 24-OCT-1997; 97US-0063045P.  
PR 24-OCT-1997; 97US-0063082P.  
PR 24-OCT-1997; 97US-0063127P.  
PR 27-OCT-1997; 97US-0063327P.  
PR 27-OCT-1997; 97US-0063329P.  
PR 28-OCT-1997; 97US-006350P.  
PR 28-OCT-1997; 97US-0063561P.  
PR 29-OCT-1997; 97US-0063704P.  
PR 29-OCT-1997; 97US-0063733P.  
PR 29-OCT-1997; 97US-0063735P.  
PR 29-OCT-1997; 97US-0063782P.  
PR 03-NOV-1997; 97US-0064288P.  
PR 07-NOV-1997; 97US-0064809P.  
PR 12-NOV-1997; 97US-0065166P.  
PR 17-NOV-1997; 97US-0065846P.  
PR 21-NOV-1997; 97US-0066364P.  
PR 24-NOV-1997; 97US-0066453P.  
PR 24-NOV-1997; 97US-0066511P.  
PR 24-NOV-1997; 97US-006670P.  
PR 11-DEC-1997; 97US-0069212P.

PR 11-DEC-1997; 97US-0069282P.  
PR 11-DEC-1997; 97US-0069334P.  
PR 16-DEC-1997; 97US-0069694P.  
PR 23-JAN-1998; 98US-0072320P.  
PR 04-FEB-1998; 98US-0073612P.  
PR 09-FEB-1998; 98US-0074086P.  
PR 09-FEB-1998; 98US-0074092P.  
PR 12-MAR-1998; 98US-0077919P.  
PR 20-MAR-1998; 98US-0078910P.  
PR 25-MAR-1998; 98US-0079294P.  
PR 27-MAR-1998; 98US-0079728P.  
PR 31-MAR-1998; 98US-0080165P.  
PR 12-JUN-1998; 98US-0081455P.  
PR 14-JUL-1998; 98US-0081455P.  
PR 28-AUG-1998; 98US-0081788P.  
PR 10-SEP-1998; 98US-0081882P.  
PR 14-SEP-1998; 98US-0081909P.  
PR 14-SEP-1998; 98US-0081909P.  
PR 14-SEP-1998; 98US-0081917P.  
PR 16-SEP-1998; 98US-0081930P.  
PR 17-SEP-1998; 98US-0081943P.  
PR 07-OCT-1998; 98US-0082114P.  
PR 29-OCT-1998; 98US-0082291P.  
PR 29-OCT-1998; 98US-0082292P.  
PR 29-OCT-1998; 98US-0082455P.  
PR 01-DEC-1998; 98US-0082510P.  
PR 05-JAN-1999; 99US-0080010P.  
PR 08-MAR-1999; 99US-0080502P.  
PR 10-MAR-1999; 99US-0080519P.  
PR 20-APR-1999; 99US-0080615P.  
PR 14-MAY-1999; 99US-0081073P.  
PR 02-JUN-1999; 99US-0081225P.  
PR 01-SEP-1999; 99US-0082011P.  
PR 08-SEP-1999; 99US-0082059P.  
PR 13-SEP-1999; 99US-0082094P.  
PR 15-SEP-1999; 99US-0082109P.  
PR 15-SEP-1999; 99US-0082154P.  
PR 05-OCT-1999; 99US-0082831P.  
PR 29-NOV-1999; 99US-0082831P.  
PR 30-NOV-1999; 99US-0082840P.  
PR 01-DEC-1999; 99US-0082830P.  
PR 01-DEC-1999; 99US-0082834P.  
PR 02-DEC-1999; 99US-0082851P.  
PR 02-DEC-1999; 99US-0082856P.  
PR 16-DEC-1999; 99US-00830095P.  
PR 20-DEC-1999; 99US-0083091P.  
PR 20-DEC-1999; 99US-0083099P.  
PR 30-DEC-1999; 99US-0083124P.  
PR 30-DEC-1999; 99US-0083127P.  
PR 05-JAN-2000; 2000US-0000219P.  
PR 06-JAN-2000; 2000US-0000277P.  
PR 06-JAN-2000; 2000US-0000317P.  
PR 11-FEB-2000; 2000US-0003365P.  
PR 18-FEB-2000; 2000US-0004341P.  
PR 22-FEB-2000; 2000US-0004342P.  
PR 24-FEB-2000; 2000US-0004414P.  
PR 24-FEB-2000; 2000US-0005004P.  
PR 01-MAR-2000; 2000US-0005601P.  
PR 02-MAR-2000; 2000US-0005746P.

(GENTH ) GENENTECH INC.  
XX  
XX Baker KP, Beresini M, DeForge L, Desnoyers L, Filvaroff E, Gao W,  
PI Gerlitsen ME, Goddard A, Godowski PJ, Gurney AL, Sherwood S,  
PI Smith V, Stewart TA, Tumas D, Watanabe CK, Wood WL, Zhang Z,  
XX WPI; 2003-352836/33.  
XX N-PSDB; ACA67255.

PT New isolated PRO polypeptide useful for treating diabetes, rheumatoid  
 PT arthritis, sports injuries, obesity, hearing loss in mammals, stroke, or  
 PT heart attack.

XX Claim 12; Fig 524; 643pp; English.

XX The present invention relates to the isolation of novel human PRO  
 CC polypeptides, and the polynucleotide sequences encoding them. The PRO  
 CC polypeptides are secreted and transmembrane proteins. The PRO  
 CC polypeptides and polynucleotides are useful for preparing a medicament  
 CC useful in the treatment of diabetes, bone and/or cartilage disorders  
 CC (e.g. rheumatoid arthritis, sports injuries, osteoarthritis), obesity,  
 CC hyper- or hypo-insulinaemia, hearing loss, and coagulation disorders  
 CC (e.g. stroke, heart attack). Anti-PRO antibodies are useful in diagnostic  
 CC assays for PRO, by detecting its expression in specific cells, tissues or  
 CC serum, and for affinity purification of PRO from recombinant cell culture  
 CC or natural sources. AB080870-AB081144 represent the human PRO  
 CC polypeptides of the invention. Note: The sequence data for this patent  
 CC was obtained in electronic format directly from the USPTO web site at  
 CC seqdata.uspto.gov/psipdb/ident.html

XX Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 6; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 6.6e-153;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALLTLVLFWGSKHFWPEVPKK 60  
 DB 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALLTLVLFWGSKHFWPEVPKK 60

QY 61 AYDMHEHTYNSGEEKKIKMEIDPVTREIFRSNGTDETLVHDFKNGYGIYFVGLOKC 120  
 DB 61 AYDMHEHTYNSGEEKKIKMEIDPVTREIFRSNGTDETLVHDFKNGYGIYFVGLOKC 120

QY 121 FIKTQIKVIFPESPEEIDEENEITTTFFEQSVIWPAPKPIENRDLKNSKILICDN 180  
 DB 121 FIKTQIKVIFPESPEEIDEENEITTTFFEQSVIWPAPKPIENRDLKNSKILICDN 180

QY 181 VTMWVNPPLISVSELDQFEEGEDLHPANKEKGIENQOMVVPQVVEKTRHARQASE 240  
 DB 181 VTMWVNPPLISVSELDQFEEGEDLHPANKEKGIENQOMVVPQVVEKTRHARQASE 240

QY 241 EELPINDYENGIEBDFPMLDERGYCCICRGNRRCRVCEPLLGYYPYPCYOGGRVYC 300  
 DB 241 EELPINDYENGIEBDFPMLDERGYCCICRGNRRCRVCEPLLGYYPYPCYOGGRVYC 300

QY 301 RVIMPCNMWVAPMLGRV 317  
 DB 301 RVIMPCNMWVAPMLGRV 317

RESULT 13

AB090933  
 ID AB090933 standard; protein; 317 AA.

XX AB090933;

DT 11-JUL-2003 (first entry)

DE Novel human secreted and transmembrane protein PRO1565.

KM Human; secreted and transmembrane protein; PRO; antibody therapy;

KM pharmaceutical; diagnostic; biosensor; bioreactor.

OS Homo sapiens.

PN US2003018173-A1.

XX 23-JAN-2003.

PF 01-MAY-2002; 2002US-00063515.

PR 06-DEC-2001; 2001US-00006867.

XX (GETH ) GENENTECH INC.

PI Batton DL, Filvaroff E, Gerritsen ME, Goddard A, Godowski PJ;

PI Grimaldi JC, Gunney AL, Katanabe CK, Wood WI,

DR WPI; 2003-401702/38.

XX N-PsDB; ACP91221.

PT New antibody useful for identifying PRO polypeptides, for affinity  
 PT purification of PRO polypeptides, and for preparing a medicament for  
 PT diagnosing or treating conditions responsive to the antibody or PRO  
 PT polypeptide.

PS Disclosure; Fig 116; 345pp; English.

XX The invention describes an antibody that specifically binds to a PRO  
 CC polypeptide having a fully defined amino acid sequence given in the  
 CC specification. The antibody is useful in identifying PRO polypeptides  
 CC useful for various industrial applications, including pharmaceuticals,  
 CC diagnostics, biosensors and bioreactors. The antibody is also used for  
 CC affinity purification of PRO polypeptides from recombinant cell culture  
 CC or natural sources. The antibody, PRO polypeptide, or its agonists or  
 CC antagonists, may be used for preparing a medicament for diagnosing or  
 CC treating a condition responsive to the antibody, PRO polypeptide, or its  
 CC agonists or antagonists. This is the amino acid sequence of a novel human  
 CC secreted and transmembrane PRO polypeptide

XX Sequence 317 AA;

Query Match 100.0%; Score 1745; DB 6; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 6.6e-153;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALLTLVLFWGSKHFWPEVPKK 60  
 DB 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALLTLVLFWGSKHFWPEVPKK 60

QY 61 AYDMHEHTYNSGEEKKIKMEIDPVTREIFRSNGTDETLVHDFKNGYGIYFVGLOKC 120  
 DB 61 AYDMHEHTYNSGEEKKIKMEIDPVTREIFRSNGTDETLVHDFKNGYGIYFVGLOKC 120

QY 121 FIKTQIKVIFPESPEEIDEENEITTTFFEQSVIWPAPKPIENRDLKNSKILICDN 180  
 DB 121 FIKTQIKVIFPESPEEIDEENEITTTFFEQSVIWPAPKPIENRDLKNSKILICDN 180

QY 181 VTMWVNPPLISVSELDQFEEGEDLHPANKEKGIENQOMVVPQVVEKTRHARQASE 240  
 DB 181 VTMWVNPPLISVSELDQFEEGEDLHPANKEKGIENQOMVVPQVVEKTRHARQASE 240

QY 241 EELPINDYENGIEBDFPMLDERGYCCICRGNRRCRVCEPLLGYYPYPCYOGGRVYC 300  
 DB 241 EELPINDYENGIEBDFPMLDERGYCCICRGNRRCRVCEPLLGYYPYPCYOGGRVYC 300

QY 301 RVIMPCNMWVAPMLGRV 317  
 DB 301 RVIMPCNMWVAPMLGRV 317

RESULT 14

AB033992  
 ID AB033992 standard; protein; 317 AA.

XX AB033992;

DT 18-SEP-2003 (first entry)

XX Human secreted/transmembrane protein PRO1565.

KM Human; secreted/transmembrane protein; PRO; tumour; cancer; cytostatic.

OS Homo sapiens.

XX US2003009013-A1.  
 XX 09-JAN-2003.  
 XX 01-MAY-2002; 2002US-00063519.  
 XX 30-DEC-1998; 98KR-00062142.  
 XX 08-MAR-1999; 99WO-US005028.  
 XX 14-MAY-1999; 99US-00311832.  
 XX 14-MAY-1999; 99WO-US010733.  
 XX 25-AUG-1999; 99US-00380137.  
 XX 25-AUG-1999; 99US-00380138.  
 XX 25-AUG-1999; 99US-00380139.  
 XX 25-AUG-1999; 99US-00380142.  
 XX 15-SEP-1999; 99US-00397342.  
 XX 18-OCT-1999; 99US-00403297.  
 XX 12-NOV-1999; 99US-00423844.  
 XX 30-DEC-1999; 99WO-US011274.  
 XX 18-FEB-2000; 2000WO-US004341.  
 XX 01-MAR-2000; 2000WO-US005601.  
 XX 02-MAR-2000; 2000WO-US005841.  
 XX 21-MAR-2000; 2000WO-US007532.  
 XX 22-MAY-2000; 2000WO-US014042.  
 XX 02-JUN-2000; 2000WO-US015264.  
 XX 22-AUG-2000; 2000WO-US023328.  
 XX 18-SEP-2000; 2000US-0064610.  
 XX 18-SEP-2000; 2000US-00665350.  
 XX 08-NOV-2000; 2000US-00709238.  
 XX 10-NOV-2000; 2000WO-US030873.  
 XX 01-DEC-2000; 2000WO-US032678.  
 XX 20-DEC-2000; 2000US-00747259.  
 XX 20-DEC-2000; 2000WO-US034956.  
 XX 28-FEB-2001; 2001WO-US006520.  
 XX 22-MAR-2001; 2001US-00816744.  
 XX 10-MAY-2001; 2001US-00854208.  
 XX 10-MAY-2001; 2001US-00854280.  
 XX 30-MAY-2001; 2001US-00870574.  
 XX 01-JUN-2001; 2001WO-US017800.  
 XX 05-JUN-2001; 2001US-00874503.  
 XX 29-JUN-2001; 2001US-00868599.  
 XX 18-JUL-2001; 2001US-00908827.  
 XX 06-DEC-2001; 2001US-00006867.  
 XX (GETH ) GENENTECH INC.  
 XX Eaton DL, Filvaroff E, Gerritsen ME, Goddard A, Godowski PJ;  
 XX Grimaldi JC, Gurney AL, Matanabe CK, Wood WI;  
 XX MPI; 2003-447384/42.  
 XX N-PDB; ACDB1598.  
 XX New isolated antibody specifically binding a PRO polypeptide, useful for  
 XX the preparation of a medicament for treating disorders with the aberrant  
 XX expression or activity of the PRO polypeptide, such as tumor conditions  
 XX and cancer.  
 XX Disclosure; Fig 116; 223pp; English.

CC present sequence represents a PRO polypeptide  
 XX Sequence 317 AA;  
 XX Query Match 100.0%; Score 1745; DB 6; Length 317;  
 XX Best Local Similarity 100.0%; Pred. No. 6,6e-153;  
 XX Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 1 MANNPENCDCIIINAEAFKSKICKSLKIGLVGIIALTLIVFMGSKHPEVPPK 60  
 DB 1 MANNPENCDCIIINAEAFKSKICKSLKIGLVGIIALTLIVFMGSKHPEVPPK 60  
 QY 61 AYMEHTFYNGEKKKIYWEIDPVRTTELFRSNGTDETLVHDFNGYTGIVGLQKC 120  
 DB 61 AYMEHTFYNGEKKKIYWEIDPVRTTELFRSNGTDETLVHDFNGYTGIVGLQKC 120  
 QY 121 FIQTQIKVPEFSEPEEEDENETTTTFEQSVIVPAKPLENDPLKSKILICDN 180  
 DB 121 FIQTQIKVPEFSEPEEEDENETTTTFEQSVIVPAKPLENDPLKSKILICDN 180  
 QY 181 VTWYINPTLISYSEIODEEEDLHFPANEKKGIEONEQWVVPQVYEKTRHAEQASE 240  
 DB 181 VTWYINPTLISYSEIODEEEDLHFPANEKKGIEONEQWVVPQVYEKTRHAEQASE 240  
 QY 241 EELPINDYENGIEFPMDERGIYCICRGKRRYCRVCEPLGTYPPYCYQGGRVIC 300  
 DB 241 EELPINDYENGIEFPMDERGIYCICRGKRRYCRVCEPLGTYPPYCYQGGRVIC 300  
 QY 301 RVIMPCNMWVYARMLGRV 317  
 DB 301 RVIMPCNMWVYARMLGRV 317  
 RESULT 15  
 ID ABU72009 standard; protein; 317 AA.  
 AC ABU72009;  
 XX 11-JUN-2003 (first entry)  
 XX Novel human secreted and transmembrane protein PRO1565.  
 XX Human; secreted and transmembrane polypeptide; chromosome mapping;  
 XX gene mapping; transgenic animal; knockout animal;  
 XX therapeutic agent screening; chromosome identification; tissue typing;  
 XX gene therapy.  
 XX Homo sapiens.  
 XX US2003018183-A1.  
 XX 23-JAN-2003.  
 XX 01-MAY-2002; 2002US-00063512.  
 XX 06-DEC-2001; 2001US-00006867.  
 XX (GETH ) GENENTECH INC.  
 XX Eaton DL, Filvaroff E, Gerritsen ME, Goddard A, Godowski PJ;  
 XX Grimaldi JC, Gurney AL, Matanabe CK, Wood WI;  
 XX MPI; 2003-330984/31.  
 XX N-PDB; ACA60420.  
 XX New secreted and transmembrane PRO polypeptides and nucleic acid  
 XX molecules encoding the polypeptides, useful in gene therapy or preparing  
 XX a medicament for treating a condition that is responsive to the PRO  
 XX polypeptide or antibody.  
 XX Disclosure; Fig 116; 409pp; English.

CC The invention describes novel isolated PRO polypeptides. The PRO  
CC polypeptides or anti-PRO antibodies are useful in preparing a medicament  
CC for treating a condition that is responsive to the PRO polypeptide or  
CC antibody. The PRO nucleotide sequences may be used as hybridization  
CC probes in chromosome and gene mapping, or in generating antisense RNA and  
CC DNA. PRO nucleic acids are also useful in preparing PRO polypeptides, in  
CC assays to identify other proteins or molecules involved in binding  
CC reaction, to generate transgenic animals or knockout animals, which in  
CC turn are useful in the development and screening of therapeutically  
CC useful reagents, for chromosome identification, and tissue typing. The  
CC PRO polypeptides and nucleic acid molecules are also useful in gene  
CC therapy, and as molecular weight markers for protein electrophoresis  
CC purposes. The anti-PRO antibodies may be used in diagnostic assays for  
CC PRO, or for the affinity purification of PRO from recombinant cell  
CC culture or natural sources. This is the amino acid sequence of a novel  
CC human secreted and transmembrane PRO polypeptide

XX Sequence 317 AA;

SQ Query Match

100.0%; Score 1745; DB 6; Length 317;

Best Local Similarity 100.0%; Pred. No. 6, 6e-153; Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKPPENCEDCHILNAEAFKSKIKCSLKIIGLVFGILALTILVFWGSKHFWPEVPKK 60  
DB 1 MAKPPENCEDCHILNAEAFKSKIKCSLKIIGLVFGILALTILVFWGSKHFWPEVPKK 60  
QY 61 AYDMEHTYSNGEKKKIYWEIDPVTREIFRSGNGTDELTLEHDFKNGYTGIVFGLQKC 120  
DB 61 AYDMEHTYSNGEKKKIYWEIDPVTREIFRSGNGTDELTLEHDFKNGYTGIVFGLQKC 120  
QY 121 FIKTQIKYIPFSEPEEEDIDNEBEITTTFFEQSVIWPAPKPIENRDFLNSKILEICDN 180  
DB 121 FIKTQIKYIPFSEPEEEDIDNEBEITTTFFEQSVIWPAPKPIENRDFLNSKILEICDN 180  
QY 181 VTMWINPTLISVSLQDFEEEGEDLHPANEKKGIENQEWVPOVKVEKTRHARQASE 240  
DB 181 VTMWINPTLISVSLQDFEEEGEDLHPANEKKGIENQEWVPOVKVEKTRHARQASE 240  
QY 241 EELPINDYTGNGIEPDPMLDERGYCCICRRGNRYCRRVCEPLLGYPPYCYOGGAVIC 300  
DB 241 EELPINDYTGNGIEPDPMLDERGYCCICRRGNRYCRRVCEPLLGYPPYCYOGGAVIC 300  
QY 301 RYIMPCNWWVARMIGRV 317  
DB 301 RYIMPCNWWVARMIGRV 317

Search completed: March 30, 2004, 14:31:03  
Job time : 62 secs



GenCore version 5.1.6  
Copyright (c) 1993 - 2004 Compugen Ltd.

## OM protein - protein search, using sw model

Run on: March 30, 2004, 14:29:47 ; Search time 22 Seconds  
(without alignments)  
743.883 Million cell updates/sec

Title: US-10-089-600-2

Perfect score: 1745

Sequence: 1 MAKNPENCEDCHILNNAE.....VICRVIMPCNMVYARMLGRV 317

Scoring table:

BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 389414 segs, 51625971 residues

Total number of hits satisfying chosen parameters: 389414

Minimum DB seq length: 0  
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%  
Maximum Match 100%

Listing first 45 summaries

Database : Issued Patents AA:  
1: /cgn2\_6/ptodata/2/1aa/5A\_COMB.pep:\*  
2: /cgn2\_6/ptodata/2/1aa/5B\_COMB.pep:\*  
3: /cgn2\_6/ptodata/2/1aa/6A\_COMB.pep:\*  
4: /cgn2\_6/ptodata/2/1aa/6B\_COMB.pep:\*  
5: /cgn2\_6/ptodata/2/1aa/PTCUS\_COMB.pep:\*  
6: /cgn2\_6/ptodata/2/1aa/backfilltest.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	521	29.9	334	1	US-08-241-465B-20
2	516	29.6	334	1	US-08-241-465B-19
3	431	24.7	296	1	US-08-241-465B-21
4	254.5	14.6	120	1	US-08-047-033-2
5	248	14.2	121	1	US-08-047-033-1
6	105.5	6.0	304	4	US-09-107-532A-738
7	97.5	5.6	374	1	US-08-464-148-2
8	97.5	5.6	374	1	US-08-385-500-2
9	97.5	5.6	374	1	US-08-846-784-2
10	92	5.3	384	4	US-09-134-001C-5242
11	92	5.3	899	4	US-09-134-000C-4697
12	91	5.2	354	3	US-09-068-569-2
13	90.5	5.2	1096	4	US-09-134-000C-5764
14	88.5	5.1	500	4	US-09-071-035-396
15	88.5	5.1	1074	4	US-09-071-035-358
16	88.5	5.1	1074	4	US-09-071-035-394
17	88	5.0	2485	5	PCT-US94-00196-1
18	87.5	5.0	1167	4	US-09-661-322A-40
19	87	5.0	1018	2	US-08-072-610-2
20	87	5.0	1018	2	US-08-719-822B-2
21	87	5.0	1018	2	US-09-092-458-2
22	86.5	5.0	368	4	US-09-328-352-7664
23	86.5	5.0	368	4	US-09-819-607-5
24	86	4.9	229	1	US-08-287-959-7
25	86	4.9	553	4	US-09-345-473E-33
26	86	4.9	2020	1	US-07-551-531-2
27	86	4.9	2818	1	US-08-510-284-1

28	86	4.9	2818	1	US-08-411-389-2	Sequence 2, Appl
29	86	4.9	2818	2	US-08-449-933-2	Sequence 2, Appl
30	86	4.9	2818	3	US-07-966-04A-2	Sequence 2, Appl
31	85.5	4.9	728	4	US-09-298-924-2	Sequence 2, Appl
32	85.5	4.9	1168	1	US-08-291-368-4	Sequence 4, Appl
33	85.5	4.9	1168	2	US-08-962-190-4	Sequence 4, Appl
34	85.5	4.9	1168	5	PCT-US95-10310-4	Sequence 4, Appl
35	85	4.9	624	4	US-09-512-563C-4	Sequence 4, Appl
36	85	4.9	854	2	US-08-926-692-17	Sequence 17, Appl
37	85	4.9	854	4	US-09-339-972-17	Sequence 17, Appl
38	85	4.9	2485	5	PCT-US94-00198-2	Sequence 2, Appl
39	84.5	4.8	369	4	US-09-819-607-2	Sequence 2, Appl
40	84.5	4.8	384	4	US-09-801-876B-6	Sequence 6, Appl
41	84.5	4.8	384	4	US-10-254-869-6	Sequence 6, Appl
42	84.5	4.8	399	4	US-09-819-607-4	Sequence 4, Appl
43	84.5	4.8	419	4	US-09-799-875-14	Sequence 14, Appl
44	84.5	4.8	700	1	US-08-726-525-7	Sequence 7, Appl
45	84.5	4.8	700	2	US-08-487-942-7	Sequence 7, Appl

## ALIGNMENTS

RESULT 1  
US-08-241-465B-20  
Sequence 20, Application US/08241465B  
Patent No. 5719125  
GENERAL INFORMATION:  
APPLICANT: Fujio SUZUKI  
APPLICANT: Yuji HIRAKI  
APPLICANT: Kazuhito TAKAHASHI  
APPLICANT: Junko SUZUKI  
APPLICANT: Jun KONDO  
APPLICANT: Atsuko KOHARA  
APPLICANT: Akiko MORI  
APPLICANT: Ei YAMADA  
TITLE OF INVENTION: HUMAN CHONDROMODULIN-I PROTEIN  
NUMBER OF SEQUENCES: 21  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: Wenderoth, Lind & Ponack  
STREET: 805 Fifteenth Street, N.W., #700  
CITY: Washington  
COUNTRY: D.C.  
ZIP: 20005  
COMPUTER READABLE FORM:  
MEDIUM TYPE: Floppy disk  
COMPUTER: IBM PC compatible  
OPERATING SYSTEM: PC-DOS/MS-DOS  
SOFTWARE: PatentIn Release #1.0, Version #1.25  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/08/241.465B  
FILING DATE: May 11, 1994  
CLASSIFICATION: 435  
ATTORNEY/AGENT INFORMATION:  
NAME: Warren M. Cheek, Jr.  
REGISTRATION NUMBER: 33,367  
REFERENCE/DOCKET NUMBER:  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (202) 371-8850  
TELEFAX:  
TELEX:  
INFORMATION FOR SEQ ID NO: 20:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 334 amino acids  
TYPE: amino acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
US-08-241-465B-20

Query Match 29.9%; Score 521; DB 1; Length 334;  
Best Local Similarity 35.5%; Pred. No. 2.9e-42;  
Matches 119; Conservative 62; Mismatches 116; Indels 38; Gaps 12;

```

Qy 6 PENCDDCHILMAEF-----SKKKICKSLKICGVFIALLTLVLFW--GSKHFWPEVP 58
Dy 15 PDDVEFC--SPPAVATLTVPSSPARLLKVGAV--LISGAVLLFGAIGAFYFMKGD 69
Qy 59 KKAADMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQ 118
Dy 70 SHIVVHYTMSINGKLQDGSMEIDAGNNLETFPKGSGAEALAVNDFONGITGIRPAGE 129
Qy 119 KCFITQIKV-IPEF-SEPEEID---ENEITTFPGQSVIWPAAKPIENRDFLKNK 173
Dy 130 KCVIAQVKARIPEVGAVTQKSISSKLEKIMPVKYEENSLIWAADQPVKONSFL-NSK 188
Qy 174 ILEICDNTWYINPTLISVSELODFEEEGDLHFPAEK-----GIEQNEQW 222
Dy 189 VLEICGDLPIFWLKPITY--PKEIQERREVRVKIVPTTKRPHSGPRSNPAGRLNNE-- 244
Qy 223 VVPOVKVETKTHARQASEEELPINDYENGIEPDPMLDERGYCCICRGNRKRCRVCEP 282
Dy 245 TRPSVQEDS-----QAFNPDPNPHYQEGESMTFPDLHDEGICICECRSRYTHCKICEP 299
Qy 283 LGGYYPYCYOGGRVICRVIMPCNMWVARMIGRV 317
Dy 300 LGGYYPMPYNYQGRSACRVIMPCSMWVARIIGMV 334

```

## RESULT 2

```

US-08-241-465B-19
; Sequence 19, Application US/08241465B
; Patent No. 5719125

```

## GENERAL INFORMATION:

```

APPLICANT: FUJIO SUZUKI
APPLICANT: YUJI HIRAKI
APPLICANT: KAZUHIRO TAKAHASHI
APPLICANT: JUNKO SUZUKI
APPLICANT: JUN KONDO
APPLICANT: ATSUKO KOHARA
APPLICANT: AKIKO MORI
APPLICANT: EI YAMADA
TITLE OF INVENTION: HUMAN CHONDROMODULIN-1 PROTEIN
NUMBER OF SEQUENCES: 21
CORRESPONDENCE ADDRESS:
ADDRESS: Wenderoth, Lind & Ponack
STREET: 805 Fifteenth Street, N.W., #700
CITY: Washington
COUNTRY: D.C.
ZIP: 20005

```

## COMPUTER READABLE FORM:

```

MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/241,465B
CLASSIFICATION: 435
FILING DATE: May 11, 1994

```

## ATTORNEY/AGENT INFORMATION:

```

NAME: Warren M. Cheek, Jr.
REGISTRATION NUMBER: 33,367
REFERENCE/DOCKET NUMBER:
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 371-8850
TELEFAX:
TELEX:

```

## INFORMATION FOR SEQ ID NO: 19:

```

SEQUENCE CHARACTERISTICS:
LENGTH: 334 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
US-08-241-465B-19

```

```

Query Match 29.6%; Score 516; DB 1; Length 334;
Best Local Similarity 35.2%; Pred. No. 8,9e-42;

```

Matches 118; Conservative 63; Mismatches 116; Indels 38; Gaps 12;

```

Qy 6 PENCDDCHILMAEF-----SKKKICKSLKICGVFIALLTLVLFW--GSKHFWPEVP 58
Dy 15 PDDVEFC--SPPAVATLTVPSSPARLLKVGAV--LISGAVLLFGAIGAFYFMKGD 69
Qy 59 KKAADMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQ 118
Dy 70 SHIVVHYTMSINGKLQDGSMEIDAGNNLETFPKGSGAEALAVNDFONGITGIRPAGE 129
Qy 119 KCFITQIKV-IPEF-SEPEEID---ENEITTFPGQSVIWPAAKPIENRDFLKNK 173
Dy 130 KCVIAQVKARIPEVGAVTQKSISSKLEKIMPVKYEENSLIWAADQPVKONSFL-SSK 188
Qy 174 ILEICDNTWYINPTLISVSELODFEEEGDLHFPAEK-----GIEQNEQW 222
Dy 189 VLEICGDLPIFWLKPITY--PKEIQERREVRVKIVPTTKRPHSGPRSNPAGRLNNE-- 244
Qy 223 VVPOVKVETKTHARQASEEELPINDYENGIEPDPMLDERGYCCICRGNRKRCRVCEP 282
Dy 245 TRPSVQEDS-----QAFNPDPNPHYQEGESMTFPDLHDEGICICECRSRYTHCKICEP 299
Qy 283 LGGYYPYCYOGGRVICRVIMPCNMWVARMIGRV 317
Dy 300 LGGYYPMPYNYQGRSACRVIMPCSMWVARIIGMV 334

```

## RESULT 3

```

US-08-241-465B-21
; Sequence 21, Application US/08241465B
; Patent No. 5719125

```

## GENERAL INFORMATION:

```

APPLICANT: FUJIO SUZUKI
APPLICANT: YUJI HIRAKI
APPLICANT: KAZUHIRO TAKAHASHI
APPLICANT: JUNKO SUZUKI
APPLICANT: JUN KONDO
APPLICANT: ATSUKO KOHARA
APPLICANT: AKIKO MORI
APPLICANT: EI YAMADA
TITLE OF INVENTION: HUMAN CHONDROMODULIN-1 PROTEIN
NUMBER OF SEQUENCES: 21
CORRESPONDENCE ADDRESS:
ADDRESS: Wenderoth, Lind & Ponack
STREET: 805 Fifteenth Street, N.W., #700
CITY: Washington
COUNTRY: D.C.
ZIP: 20005

```

## COMPUTER READABLE FORM:

```

MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/241,465B
CLASSIFICATION: 435
FILING DATE: May 11, 1994

```

## ATTORNEY/AGENT INFORMATION:

```

NAME: Warren M. Cheek, Jr.
REGISTRATION NUMBER: 33,367
REFERENCE/DOCKET NUMBER:
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 371-8850
TELEFAX:
TELEX:

```

## INFORMATION FOR SEQ ID NO: 21:

```

SEQUENCE CHARACTERISTICS:
LENGTH: 296 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
US-08-241-465B-21

```

Query Match 24.7% Score 431; DB 1; Length 296;  
Best Local Similarity 31.2% Pred. No. 1.3e-33;  
Matches 103; Conservative 57; Mismatches 104; Indels 66; Gaps 11;

DB 6 PENCDCCHILNARF-----KSKKICKSLKICGLVFGIALTLVLFW--GSKHWPFPV 58  
15 PDVDFEC--SPPAVATLVKSPSPARLLKGAVV--LISGAVLLFGAIGAFYFWKSD 69  
QY 59 KRAYDMHETFGSGEKKIYMEIDPVTREIFRSNGTDELEVDENKNGYIGITFVGQ 118  
70 SHIYVHYTMSINGKLQDGSMEIDAGNNLETFMGSGAEALAVDPQ----- 118  
QY 119 KCFIKQIIVIPFSPSEEDIDENEITTFPEOSYIWPAPKPLENRDPLNKLILIC 178  
119 -----EGKMP-----VKYENSLLIWAADQPVKDNFL--SKVLELC 155  
QY 179 DNVTMWINPTLISVELQDFEEGEDLHPANEK-----GIEQNEQVNVQY 227  
156 GDLPIFMLKPTY--PREIQERRREVVKIVPTTKRPHNGPSPNPGAGRLNNE--TRBSV 211  
QY 228 KVEKTHARQASEELPINDYTENGIEFDPMLDERGYCCYCRGRNRYCRVCEPLIGY 287  
212 QEDS-----QAFPDNPHYQEGESMTFPRLDHBCICIEGRSYTHCQKICEPLGGY 266  
QY 288 PYPYCYQGRVIGRVIMPCNMWVARMGRV 317  
DB 267 PMPYNYQGRSACRVIMPCSMWVARILGMV 296

RESULT 4  
US-08-047-033-2  
; Sequence 2, Application US/08047033  
; Patent No. 5444157  
; GENERAL INFORMATION:  
; APPLICANT: Fujio SUZUKI et al.  
; TITLE OF INVENTION: CHONDROMODULIN-I PROTEIN  
; NUMBER OF SEQUENCES: 12  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Wenderoth, Lind & Ponack  
; STREET: 805 Fifteenth Street, N.W., #700  
; CITY: Washington  
; STATE: D.C.  
; COUNTRY: U.S.A.  
; ZIP: 20005  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: Diskette, 5.25 inch, 500 Kb  
; COMPUTER: IBM Compatible  
; OPERATING SYSTEM: MS-DOS  
; SOFTWARE: DisplayWrite  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/047,033  
; FILING DATE:  
; CLASSIFICATION: 514  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: US/07/821,859  
; FILING DATE:  
; APPLICATION NUMBER: US/07/745,497  
; FILING DATE:  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Warren M. Cheek Jr.  
; REGISTRATION NUMBER: 33,367  
; REFERENCE/DOCKET NUMBER:  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: 202-371-8850  
; TELEFAX: 202-371-8856  
; TELEX:  
; INFORMATION FOR SEQ ID NO: 2:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 120 amino acids  
; TYPE: amino acid  
; STRANDEDNESS: single  
; TOPOLOGY: linear  
; MOLECULE TYPE: protein

HYPOTHETICAL:  
ANTI-SENSE:  
FRAGMENT TYPE:  
ORIGINAL SOURCE:  
ORGANISM: bovine  
STRAIN:  
INDIVIDUAL ISOLATE:  
DEVELOPMENTAL STAGE:  
HAPLOTYPE:  
TISSUE TYPE: fetal cartilage  
CELL LINE:  
ORGANELLE:  
IMMEDIATE SOURCE:  
LIBRARY:  
CLONE:  
POSITION IN GENOME:  
CHROMOSOME/SEGMENT:  
MAP POSITION:  
UNITS:  
FEATURE:  
NAME/KEY: modified site  
LOCATION: 7  
IDENTIFICATION METHOD:  
OTHER INFORMATION: /note= "Xaa is Met or Val"  
FEATURE:  
NAME/KEY: modified site  
LOCATION: 10  
IDENTIFICATION METHOD:  
OTHER INFORMATION: /note= "Xaa is Glu or Thr"  
FEATURE:  
NAME/KEY: modified site  
LOCATION: 83  
IDENTIFICATION METHOD:  
OTHER INFORMATION: /note= "Xaa is Cys or Val"  
PUBLICATION INFORMATION:  
AUTHORS:  
TITLE:  
JOURNAL:  
VOLUME:  
ISSUE:  
PAGES:  
DATE:  
DOCUMENT NUMBER:  
FILING DATE:  
PUBLICATION DATE:  
RELEVANT RESIDUES IN SEQ ID NO:  
US-08-047-033-2

Query Match 14.6% Score 254.5; DB 1; Length 120;  
Best Local Similarity 53.9% Pred. No. 4.3e-17;  
Matches 48; Conservative 11; Mismatches 27; Indels 3; Gaps 2;

QY 232 TRHARQASEELP-PINDYTENG--IEFDPMLDERGYCCYCRGRNRYCRVCEPLIGY 288  
DB 32 TRPSVQDAEPFNPDPHYQGESMTFPRLDHBCICIEGRSYTHCQKICEPLGGY 91  
QY 289 YPYCYQGRVIGRVIMPCNMWVARMGRV 317  
DB 92 WPMYNYQGRSACRVIMPCSMWVARILGMV 120

RESULT 5  
US-08-047-033-1  
; Sequence 1, Application US/08047033  
; Patent No. 5444157  
; GENERAL INFORMATION:  
; APPLICANT: Fujio SUZUKI et al.  
; TITLE OF INVENTION: CHONDROMODULIN-I PROTEIN  
; NUMBER OF SEQUENCES: 12  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Wenderoth, Lind & Ponack  
; STREET: 805 Fifteenth Street, N.W., #700

CITY: Washington  
STATE: D.C.  
COUNTRY: U.S.A.  
ZIP: 20005  
COMPUTER READABLE FORM:  
MEDIUM TYPE: Diskette, 5.25 inch, 500 Kb  
COMPUTER: IBM compatible  
OPERATING SYSTEM: MS-DOS  
SOFTWARE: DisplayWrite  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/08/047,033  
FILING DATE:  
CLASSIFICATION: 514  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: US/07/821,859  
FILING DATE:  
APPLICATION NUMBER: US/07/745,497  
FILING DATE:  
ATTORNEY/AGENT INFORMATION:  
NAME: Warren M. Cheek, Jr.  
REGISTRATION NUMBER: 33,367  
REFERENCE/DOCKET NUMBER:  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: 202-371-8850  
TELEFAX: 202-371-8856  
TELEX:  
INFORMATION FOR SEQ. ID NO: 1:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 121 amino acids  
TYPE: amino acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
MOLECULE TYPE: protein  
HYPOTHETICAL:  
ANTI-SENSE:  
FRAGMENT TYPE:  
ORIGINAL SOURCE:  
ORGANISM: bovine  
STRAIN:  
INDIVIDUAL ISOLATE:  
DEVELOPMENTAL STAGE:  
HAPLOTYPE:  
TISSUE TYPE: fetal cartilage  
CELL TYPE:  
CELL LINE:  
ORGANELLE:  
IMMEDIATE SOURCE:  
LIBRARY:  
CLONE:  
POSITION IN GENOME:  
CHROMOSOME/SEGMENT:  
MAP POSITION:  
UNITS:  
FEATURE:  
NAME/KEY: modified site  
LOCATION: 7  
IDENTIFICATION METHOD:  
OTHER INFORMATION: /note= "Xaa is Met or Val"  
FEATURE:  
NAME/KEY: modified site  
LOCATION: 10  
IDENTIFICATION METHOD:  
OTHER INFORMATION: /note= "Xaa is Glu or Thr"  
FEATURE:  
NAME/KEY: modified site  
LOCATION: 84  
IDENTIFICATION METHOD:  
OTHER INFORMATION: /note= "Xaa is Cys or Val"  
PUBLICATION INFORMATION:  
AUTHORS:  
TITLE:  
JOURNAL:  
VOLUME:

ISSUE:  
PAGES:  
DATE:  
DOCUMENT NUMBER:  
FILING DATE:  
PUBLICATION DATE:  
RELEVANT RESIDUES IN SEQ ID NO:  
US-08-047-033-1

Query Match 14.2%; Score 248; DB 1; Length 121;  
Best Local Similarity 52.2%; Pred. No. 1.9e-16;  
Matches 47; Conservative 11; Mismatches 28; Indels 4; Gaps 2;

Qy 232 TRHARQASEEL-PINDYTE--NGIEFDMLERGYCCICRRGNRCRVCPEPLIGY 287  
Db 32 TRFSVQDAEPFNDPNFHQEGESMTFDPDLDEGICIECRSRSTHCKIXEPLOGYH 91  
Qy 288 PYPYCYOGGRVYICRVIMPCNMWVARMUGRV 317  
Db 92 PWPYNYGCRSACRVIMPCSMWVARILIGVY 121

RESULT 6  
US-09-107-532A-7138  
Sequence 7138, Application US/09107532A  
Patent No. 6583275  
GENERAL INFORMATION:  
APPLICANT: Lynn A. Doucette-Stamm and David Bush  
TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO  
ENTEROCOCCUS FAECIUM FOR DIAGNOSTICS AND THERAPEUTICS  
NUMBER OF SEQUENCES: 7310  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: GENOME THERAPEUTICS CORPORATION  
STREET: 100 Beaver Street  
CITY: Walpole  
STATE: Massachusetts  
COUNTRY: USA  
ZIP: 02354  
COMPUTER READABLE FORM:  
MEDIUM TYPE: CD-ROM ISO9660  
COMPUTER: PC  
OPERATING SYSTEM: <Unknown>  
SOFTWARE: ASCII  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/09/107,532A  
FILING DATE: 30-Jun-1998  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: 60/085,598  
FILING DATE: 14 May 1998  
APPLICATION NUMBER: 60/051571  
FILING DATE: July 2, 1997  
ATTORNEY/AGENT INFORMATION:  
NAME: Atinello, Pamela Deneke  
REGISTRATION NUMBER: 40,489  
REFERENCE/DOCKET NUMBER: GTC-012  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (781)893-5007  
TELEFAX: (781)893-8277  
INFORMATION FOR SEQ ID NO: 7138:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 304 amino acids  
TYPE: amino acid  
TOPOLOGY: linear  
MOLECULE TYPE: protein  
HYPOTHETICAL: YES  
ORIGINAL SOURCE:  
ORGANISM: Enterococcus faecium  
FEATURE:  
NAME/KEY: misc feature  
LOCATION: (B) LOCATION 1...304  
SEQUENCE DESCRIPTION: SEQ ID NO: 7138:  
US-09-107-532A-7138

Query Match 6.0%; Score 105.5; DB 4; Length 304;  
 Best Local Similarity 19.5%; Pred. No. 0.043;  
 Matches 55; Conservative 48; Mismatches 104; Indels 75; Gaps 9;

QY 17 AEAERKSKICKSLKI-CGLVFGILALTLI-VLWFSKHFWEVPAKADMEHTFYSNGEK-74  
 DB 25 AQIDKVKYKTKGVLTYGALIGFLAIGMILV-----FSPYVMGKSYDFETVXKSSAQN-78  
 QY 75 -KXIYMEIDPVT-----RTEIFRSGNGTDETLVH-103  
 DB 79 TSSIANMYDDWEINREAGLEFLVLRYPQDDTGSLSNIKSYKLNFTNKNNEDVKTQV-138  
 QY 104 DFNKGYTGIVFGLOKCFIKTQIKVIRPF-----SEPEEIDENE-----143  
 DB 139 KLSDEYTYIVYQNLPKDGVISATVNPRIYYPELEPNSDKKEIKFYAVDKDIXNAKL-158  
 QY 144 EITTFPQSVIWPVPAEKPIEN--RDFIKNSKILIEDNTVMWINPILSVSELDPFEE-201  
 DB 199 EVESTSLQKDNFYQIQAILENSIRDEKKSIEKNLAKVQ-----KQIDK-245

QY 202 EGEDLHFPANERKGIQNEQWVVPQVKEKTRHARQASEEEL-243  
 DB 246 ANQNLFPOTLDEQDDTKNEISSM-QTTENNKSSIESEENKI-286

RESULT 7  
 US-08-464-148-2  
 ; Sequence 2, Application US/08464148  
 ; Patent No. 5710026  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Sprecher, Cindy A.  
 ; TITLE OF INVENTION: CYTOPLASMIC ANTIPROTEINASE-2 AND  
 ; TITLE OF INVENTION: CYTOPLASMIC ANTIPROTEINASE-3 AND CODING SEQUENCES  
 ; NUMBER OF SEQUENCES: 16  
 ; CORRESPONDENCE ADDRESS:  
 ; ADDRESSEE: Townsend and Townsend Kourie and Crew  
 ; STREET: Stuart Street Tower, One Market Plaza  
 ; CITY: San Francisco  
 ; STATE: California  
 ; COUNTRY: US  
 ; ZIP: 94105-1493  
 ; COMPUTER READABLE FORM:  
 ; MEDIUM TYPE: Floppy disk  
 ; COMPUTER: IBM PC compatible  
 ; OPERATING SYSTEM: PC-DOS/MS-DOS  
 ; SOFTWARE: Patentin Release #1.0, Version #1.25  
 ; CURRENT APPLICATION DATA:  
 ; APPLICATION NUMBER: US/08/464,148  
 ; FILING DATE: 05-JUN-1995  
 ; CLASSIFICATION: 435  
 ; PRIORITY APPLICATION DATA:  
 ; APPLICATION NUMBER: US 08/385,500  
 ; FILING DATE: 08-FEB-1995  
 ; ATTORNEY/AGENT INFORMATION:  
 ; NAME: Parmelee, Steven W.  
 ; REGISTRATION NUMBER: 31,990  
 ; REFERENCE/DOCKET NUMBER: 13952-21  
 ; TELECOMMUNICATION INFORMATION:  
 ; TELEPHONE: (206) 467-9600  
 ; TELEFAX: (415) 543-5043  
 ; INFORMATION FOR SEQ ID NO: 2:  
 ; SEQUENCE CHARACTERISTICS:  
 ; LENGTH: 374 amino acids  
 ; TYPE: amino acid  
 ; TOPOLOGY: linear  
 ; MOLECULE TYPE: protein  
 ; US-08-464-148-2

Query Match 5.6%; Score 97.5; DB 1; Length 374;  
 Best Local Similarity 21.3%; Pred. No. 0.35;  
 Matches 63; Conservative 42; Mismatches 112; Indels 79; Gaps 14;

QY 7 ENCEDC--HINAEAFKSKICKSLKICGLVFGILALTLI--VLWFSKHFWEVPAKAY-62

DB 123 EDTTECRKHINQWVAEKTEGKISEVLDAGTVDPITKVLVNAIYFGK--WNBOFDRKY-179  
 QY 63 DMEHTFYSNGEKKKIYMEIDPVTREIFRSGNGTDETLVHDFKNGYTGIVFGLOKCFI-122  
 DB 180 TRGMLFKTNEKKTYQM-----WFKKAK-----FQNGYAD-----EV-211  
 QY 123 KIQIKVIRPFSEPEEEL-----DENEIITT-----FPGSVIWPVPAEKPIENR--DFL-169  
 DB 212 HQVLELPYV--EELSVMILLPDDNTDLAVVEKALTYEKKRAMTNSKELTKSKQVFL-268  
 QY 170 KNSKILIEDNTVMWINPILSVSELQPEEEDLHFPANERK-----GIBON-219  
 DB 269 PRKLEE-----SYDLEPFLRLGNTDAFDKADFGSGMSTKXVPLSKVAHKCFEVRN-322  
 QY 220 EQWVVPQVKEKTRHARQASEEELPINDYENGIEFDPML-----DERGYCCYICRR-271  
 DB 323 EEGTEAANAATAVVRNRCGRMEPRFCADH-----PFLFTRRHKTNCILFCGR-370

RESULT 8  
 US-08-385-500-2  
 ; Sequence 2, Application US/08385500  
 ; Patent No. 5712117  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Sprecher, Cindy A.  
 ; TITLE OF INVENTION: CYTOPLASMIC ANTIPROTEINASE-2 AND  
 ; TITLE OF INVENTION: CYTOPLASMIC ANTIPROTEINASE-3 AND CODING SEQUENCES  
 ; NUMBER OF SEQUENCES: 16  
 ; CORRESPONDENCE ADDRESS:  
 ; ADDRESSEE: Townsend and Townsend Kourie and Crew  
 ; STREET: Stuart Street Tower, One Market Plaza  
 ; CITY: San Francisco  
 ; STATE: California  
 ; COUNTRY: US  
 ; ZIP: 94105-1493  
 ; COMPUTER READABLE FORM:  
 ; MEDIUM TYPE: Floppy disk  
 ; COMPUTER: IBM PC compatible  
 ; OPERATING SYSTEM: PC-DOS/MS-DOS  
 ; SOFTWARE: Patentin Release #1.0, Version #1.25  
 ; CURRENT APPLICATION DATA:  
 ; APPLICATION NUMBER: US/08/385,500  
 ; FILING DATE:  
 ; CLASSIFICATION: 435  
 ; ATTORNEY/AGENT INFORMATION:  
 ; NAME: Parmelee, Steven W.  
 ; REGISTRATION NUMBER: 31,990  
 ; REFERENCE/DOCKET NUMBER: 13952-21  
 ; TELECOMMUNICATION INFORMATION:  
 ; TELEPHONE: (206) 467-9600  
 ; TELEFAX: (415) 543-5043  
 ; INFORMATION FOR SEQ ID NO: 2:  
 ; SEQUENCE CHARACTERISTICS:  
 ; LENGTH: 374 amino acids  
 ; TYPE: amino acid  
 ; TOPOLOGY: linear  
 ; MOLECULE TYPE: protein  
 ; US-08-385-500-2

Query Match 5.6%; Score 97.5; DB 1; Length 374;  
 Best Local Similarity 21.3%; Pred. No. 0.35;  
 Matches 63; Conservative 42; Mismatches 112; Indels 79; Gaps 14;

QY 7 ENCEDC--HINAEAFKSKICKSLKICGLVFGILALTLI--VLWFSKHFWEVPAKAY-62  
 DB 123 EDTTECRKHINQWVAEKTEGKISEVLDAGTVDPITKVLVNAIYFGK--WNBOFDRKY-179  
 QY 63 DMEHTFYSNGEKKKIYMEIDPVTREIFRSGNGTDETLVHDFKNGYTGIVFGLOKCFI-122  
 DB 180 TRGMLFKTNEKKTYQM-----WFKKAK-----FQNGYAD-----EV-211  
 QY 123 KIQIKVIRPFSEPEEEL-----DENEIITT-----FPGSVIWPVPAEKPIENR--DFL-169

Db 212 HTQVLELPYV---EELISWYLLPDNDTDLAVVEKALTYEKFKAMTNSKLTFSKVQVFL 268  
 QY 170 KSKKILEICDNVTMYINIPFLISVSELOPFEEGEDLHPANCK-----GIEON 219  
 Db 269 PRKLEB-----SYDLEPFLRLGMDADAEKADPSGNSTEKNVPLSKVAHKCFVEVN 322  
 QY 220 EQWVVPQVYKVKTRHARQASEELPINDYENGIEPDPML---DERGYCCICYCR 271  
 Db 323 EEGTEAATAATVAVNRSRCSMRERFCADH-----PFLFIRRHKTNCILFCGR 370

## RESULT 9

US-08-846-784-2  
 ; Sequence 2, Application US/08846784  
 ; Patent No. 5747645  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Sprecher, Cindy A.  
 ; TITLE OF INVENTION: CYTOSOLIC ANTIPROTEINASE-2 AND  
 ; TITLE OF INVENTION: CYTOSOLIC ANTIPROTEINASE-3 AND CODING SEQUENCES  
 ; NUMBER OF SEQUENCES: 16  
 ; CORRESPONDENCE ADDRESS:  
 ; ADDRESSEE: Townsend and Townsend Khourie and Crew  
 ; STREET: Stewart Street Tower, One Market Plaza  
 ; CITY: San Francisco  
 ; STATE: California  
 ; COUNTRY: US  
 ; ZIP: 94105-1493  
 ; COMPUTER READABLE FORM:  
 ; MEDIUM TYPE: Floppy disk  
 ; COMPUTER: IBM PC compatible  
 ; OPERATING SYSTEM: PC-DOS/MS-DOS  
 ; SOFTWARE: Patent Release #1.0, Version #1.25  
 ; CURRENT APPLICATION DATA:  
 ; APPLICATION NUMBER: US/08/846,784  
 ; FILING DATE: 30-APR-1997  
 ; CLASSIFICATION: 435  
 ; PRIOR APPLICATION DATA:  
 ; APPLICATION NUMBER: 08/385,500  
 ; FILING DATE: 08-FEB-1995  
 ; ATTORNEY/AGENT INFORMATION:  
 ; NAME: Parmelee, Steven W.  
 ; REGISTRATION NUMBER: 31,990  
 ; REFERENCE/DOCKET NUMBER: 13952-21  
 ; TELECOMMUNICATION INFORMATION:  
 ; TELEPHONE: (206) 467-9600  
 ; TELEFAX: (415) 543-5043  
 ; INFORMATION FOR SEQ ID NO: 2:  
 ; SEQUENCE CHARACTERISTICS:  
 ; LENGTH: 374 amino acids  
 ; TYPE: amino acid  
 ; TOPOLOGY: linear  
 ; MOLECULE TYPE: protein  
 ; US-08-846-784-2

Query Match 5.6%; Score 97.5; DB 1; Length 374;  
 Best Local Similarity 21.3%; Pred. No. 0.35;  
 Matches 63; Conservative 42; Mismatches 112; Indels 79; Gaps 14;

QY 7 ENCEDC--HINAFAFSKKIKCKSLKCGLVFGILATLL--VLFMSKHFMEVPEKAY 62  
 Db 123 EDTECKRHINDVAEKTGKISVLDAGYDPLTKLVNVAIFYKSK--WNEQDRKY 179  
 QY 63 DMEHTFYSNGEKKIYWEIDPVTREIFRSNGTDETLVHDPFGNGYTGIVYGLQKCFI 122  
 Db 180 TRGMLFKTNEKKITVQM-----MFKEAK-----FMGYAD-----EV 211  
 QY 123 KTQIKVPERSEPEEL-----DENEITTT-----FFEOSVIVAEKPIENR--DFI 169  
 Db 212 HTQVLELPYV---EELISWYLLPDNDTDLAVVEKALTYEKFKAMTNSKLTFSKVQVFL 268  
 QY 170 KSKKILEICDNVTMYINIPFLISVSELOPFEEGEDLHPANCK-----GIEON 219

Db 269 PRKLEB-----SYDLEPFLRLGMDADAEKADPSGNSTEKNVPLSKVAHKCFVEVN 322  
 QY 220 EQWVVPQVYKVKTRHARQASEELPINDYENGIEPDPML---DERGYCCICYCR 271  
 Db 323 EEGTEAATAATVAVNRSRCSMRERFCADH-----PFLFIRRHKTNCILFCGR 370

## RESULT 10

US-09-134-001C-5242  
 ; Sequence 5242, Application US/09134001C  
 ; Patent No. 6380370  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Lynn Doucette-Stamm et al  
 ; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO STAPHYLOCOCCUS  
 ; TITLE OF INVENTION: EPIDERMIDIS FOR DIAGNOSTICS AND THERAPEUTICS  
 ; FILE REFERENCE: GTC-007  
 ; CURRENT APPLICATION NUMBER: US/09/134,001C  
 ; PRIOR FILING DATE: 1997-11-08  
 ; PRIOR APPLICATION NUMBER: US 60/064,964  
 ; PRIOR FILING DATE: 1997-08-14  
 ; NUMBER OF SEQ ID NOS: 5674  
 ; SEQ ID NO 5242  
 ; LENGTH: 384  
 ; TYPE: PRT  
 ; ORGANISM: Staphylococcus epidermidis  
 ; US-09-134-001C-5242

Query Match 5.3%; Score 92; DB 4; Length 384;  
 Best Local Similarity 26.4%; Pred. No. 1.2;  
 Matches 56; Conservative 36; Mismatches 62; Indels 58; Gaps 13;

QY 88 EIFRSGNTDTELVHDPKNG---YTGIFYVGLQKCFIKQIKYIPERSSEEDISNEE 144  
 Db 145 EIHHDGN-----ITYONKNGGIPATGLVKTGKTK--KGTQVY--FKPDSIFKS-- 191  
 QY 145 ITTFP-FEOSVIVPAEKPIENRDLNNSKILICDNVTMYINIPFLISVSELOPFEEG 203  
 Db 192 -TTTFNFD-----ILSERLOESAFILKDLK-----ITLTDLRSKERE 228  
 QY 204 EDLHPANCKGIEONEGV--VPOYKVKTRHARQASEELPI-----ND-YTENGIEF- 255  
 Db 229 EIVHY---EGIKRFPYSVYNGEKEVLHVDVTFGHSNGLEVDVAFQINDYSESILSFV 284  
 QY 256 -----DPMLEDERGYCCICYCRGNRYCRRVCE 281  
 Db 285 NNVRTKDGTHGVGFKTAMTRVFNEVARRINE 316

RESULT 11  
 US-09-134-000C-4697  
 ; Sequence 4697, Application US/09134000C  
 ; Patent No. 6617156  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Lynn Doucette-Stamm et al  
 ; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO  
 ; TITLE OF INVENTION: ENTEROCOCCUS FAECALIS FOR DIAGNOSTICS AND THERAPEUTICS  
 ; FILE REFERENCE: 032796-032  
 ; CURRENT APPLICATION NUMBER: US/09/134,000C  
 ; PRIOR FILING DATE: 1998-08-13  
 ; PRIOR APPLICATION NUMBER: US 60/055,778  
 ; PRIOR FILING DATE: 1997-08-15  
 ; NUMBER OF SEQ ID NOS: 6812  
 ; SOFTWARE: Patent version 3.1  
 ; SEQ ID NO 4697  
 ; LENGTH: 899  
 ; TYPE: PRT  
 ; ORGANISM: Enterococcus faecalis  
 ; US-09-134-000C-4697

Query Match 5.3%; Score 92; DB 4; Length 899;  
 Best Local Similarity 19.2%; Pred. No. 4.1;

Matches 53; Conservative 40; Mismatches 99; Indels 84; Gaps 12;  
QY 68 FYSNGEKKI-----YMEIDPTREIFRSNGTDELEVDHDFKNGYIVFGQKQFIK 123  
DB 461 FTTPKXVRLSDSDYFADPFKORLSKVSQCI--IVEVEDLATYSGIFRLTKAKYLT 518  
QY 124 TQIKVIDEF-----SEPEEIDENE-----ITTFEPQSV 154  
DB 519 ANKIVAEQYKLEDTYTSNQGVIITRNEDTYKDVFEFTKANKVSAGVLTAKTAEKTK 578  
QY 155 IWVAEKEIENRDLKSKILEICDNTMWI-----NPTLISVSELOPFEEDGELHF 208  
DB 579 SGIRLQKANGYITANKNVV---ATGSWIANHTVNEQIIMKNSDNFYGSDPLY- 633  
QY 209 PANEGKIEQNEQWVQVKEKTRHARQASEELPINDYTENGIEFDPMLDERGYCCIIY 268  
DB 634 -----KGAASKSGSLVPVGI-----YREN--QVRLITONG-----Y 665  
QY 269 CRGRNRYGRVCEPLG--YYP-----YCYQ 294  
DB 666 LTANKSYAQKVPNIKDYLYPEYVWKTNDYYQ 701

## RESULT 12

US-09-068-569-2  
; Sequence 2, Application US/09068569  
; Patent No. 6238859  
; GENERAL INFORMATION:  
; APPLICANT: Like, Wolfgang  
; APPLICANT: Hunsmann, Gerhard  
; APPLICANT: Weber, Thomas  
; TITLE OF INVENTION: Virus Protein Antigens of the JC Virus  
; FILE REFERENCE: Like et al  
; CURRENT APPLICATION NUMBER: US/09/068,569  
; CURRENT FILING DATE: 1998-07-17  
; EARLIER APPLICATION NUMBER: PCT/EP96/05177  
; EARLIER FILING DATE: 1996-11-22  
; EARLIER APPLICATION NUMBER: DE 195 43 553.2  
; EARLIER FILING DATE: 1995-11-22  
; NUMBER OF SEQ ID NOS: 4  
; SOFTWARE: PatentIn Ver. 2.0  
; SEQ ID NO 2  
; LENGTH: 354  
; TYPE: PRT  
; ORGANISM: JC virus  
US-09-068-569-2

Query Match 5.2%; Score 91; DB 3; Length 354;  
Best Local Similarity 21.5%; Pred. No. 1.4;  
Matches 61; Conservative 27; Mismatches 86; Indels 110; Gaps 15;

QY 72 GKKKIWEIDPVRTIEFRSGNGTDELEVDHDFKNGYIVFGQKQFIKQIKYIPE 131  
DB 8 GERK-----DPVOYKILIRG-----VEVLEKVTGVDSTEV---ECFL-----TPE 47  
QY 132 FSEPEEEL--DENEITTTTFEQSVIWPAPKEIENRDLKSKILEI-----C 178  
DB 48 MGDDDEHLRFSKSIISDTF-----ESDSPRDMPCYSVARILPLINLBDLTC 97  
QY 179 DNVTMWTNPTL-----ISVSELOPFEEDGELH----- 207  
DB 98 GNILM-WEAVTLKTEVIGVSLKMWVHNSGQTHUNGAGKPYQSTSPFFSVGGEALDQ 156  
QY 208 -----FPAN-----EKKGIEQNE-----EQWVVPQVKEKTRH--A 235  
DB 157 VLFNRYTKYPDGTLFPKNATVQSQVNTENKAYLDKAKAVVEGWDVPRNENTRYFGT 216  
QY 236 RQASEEELPINDYTENGIEFDPMLDERGYCCIIYCRGRNRYGRV 279  
DB 217 LTGENVPEVLAHTNTATTV--LLDEFGVGPL-CKGDNLYLSAV 257

## RESULT 13

US-09-134-000C-5764  
; Sequence 5764, Application US/09134000C  
; Patent No. 6617156  
; GENERAL INFORMATION:  
; APPLICANT: Lynn Doucette-Stamm et al  
; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO  
; TITLE OF INVENTION: ENTEROCOCCUS FAECALIS FOR DIAGNOSTICS AND THERAPEUTICS  
; FILE REFERENCE: 032796-032  
; CURRENT APPLICATION NUMBER: US/09/134,000C  
; CURRENT FILING DATE: 1998-08-13  
; PRIOR APPLICATION NUMBER: US 60/055,778  
; PRIOR FILING DATE: 1997-08-15  
; NUMBER OF SEQ ID NOS: 6812  
; SOFTWARE: PatentIn version 3.1  
; SEQ ID NO 5764  
; LENGTH: 1096  
; TYPE: PRT  
; ORGANISM: Enterococcus faecalis  
US-09-134-000C-5764

Query Match 5.2%; Score 90.5; DB 4; Length 1096;  
Best Local Similarity 19.3%; Pred. No. 7.7;  
Matches 53; Conservative 41; Mismatches 86; Indels 95; Gaps 12;

QY 50 SKHFMPEV-PKXAYMEHTFY-----SNGEKKIWEIDPVRTIEFRSG----- 93  
DB 577 NEFMQETAPRPGYTLDETXYVSIKYDNNKNAVTR-DYTAKEQVIRFGDFPKFAG 635  
QY 94 -----NGTDELEVDH-----FKNGYTIYFV- 115  
DB 636 SADGTAETGFNDLSPKVSPLSGTNEITGAEDKATACNBOGLGFDGYGKFNLPYGDYLLE 695  
QY 116 -----GLQCKYKIQIKYIPFSPEEIDENEITTFEQ-----SVIWPAPKEPIE 164  
DB 696 EIEAPGFGQKI--PPLERSTFKENKODYAKSEYVFTTTEGQROPIMVTPYEK-- 749  
QY 165 NBDPLKSKILEICDNTMWIWNPTLISVSELO-----DPEEEDGELHFPANE 212  
DB 750 -----LTNNEFSVSLRMIMYDPEKEDSLTSLATWKDGKKLNTLDFELVAKLNYNHE 805  
QY 213 KKGIEQNEQW--VPQVKEKTRHARQASEELPI 245  
DB 806 IK-----EDWYVVAQADIVEATKAQOEKDEKAPV 835

## RESULT 14

US-09-071-035-396  
; Sequence 396, Application US/09071035  
; Patent No. 6448043  
; GENERAL INFORMATION:  
; APPLICANT: Gil H. Choi  
; TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides  
; NUMBER OF SEQUENCES: 496  
; CORRESPONDENCE ADDRESS:  
; ADDRESSER: Human Genome Sciences, Inc.  
; STREET: 9410 Key West Avenue  
; CITY: Rockville  
; STATE: Maryland  
; COUNTRY: USA  
; ZIP: 20850  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage  
; COMPUTER: HP Vectra 486/33  
; OPERATING SYSTEM: MSDOS version 6.2  
; SOFTWARE: ASCII Text  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/09/071,035  
; FILING DATE:  
; CLASSIFICATION:  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER:  
; FILING DATE:  
; ATTORNEY/AGENT INFORMATION:

NAME: A. Anders Brookes  
REGISTRATION NUMBER: 36,373  
REFERENCE/DOCKET NUMBER: PB369P2  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (301) 309-8504  
TELEFAX: (301) 309-8512  
INFORMATION FOR SEQ ID NO: 396:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 500 amino acids  
TYPE: amino acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
MOLECULE TYPE: protein  
US-09-071-035-356

Query Match 5.1%; Score 88.5; DB 4; Length 500;  
Best Local Similarity 19.3%; Pred. No. 3.9;  
Matches 53; Conservative 40; Mismatches 87; Indels 95; Gaps 12;

QY 50 SKHFWPEV-PKXAYDMEHTFY-----SNGKKKIYMEIDVTRTEIFRSG----- 93  
DB 15 NEYFWOETKAPESYTLDETKYPSIKVDNNEKNVITR-DYTAKEQVIRFGDFPKFAG 73  
QY 94 -----NGTDETLFVHD-----FKNGYTGIFYV- 115  
DB 74 SADGTATGFNDLSEKVSPLSGTIXEITGADKATTACNEQLGPDGKFNLEPGDYLL 133  
QY 116 -----GLQKCFIKTQIKVIPSEPEEIDENEETITTFEQ-----SVIWPAPKPIE 164  
DB 134 EIEAPBGFQKI--TPLEIRSTFKENKDYAKSEVYFTITEBQKQPIKMTVPYK-- 187  
QY 165 NRDFLXSKILTEICDVTWYINPTLISVSELQ-----DFEEEGDLHFPPANE 212  
DB 188 ----LNNPFSVSLNRLMYDLPEKEDSLSLATWMDGKNKNTLDFTLVDLRLNLS 243  
QY 213 KKGIEQNEQW--VVPQYKVEKTRHARQASEELPI 245  
DB 244 IK-----EDWYVVAQAIDVEATKAAQEKDEKAPV 273

## RESULT 15

US-09-071-035-356  
Sequence 358, Application US/09071035  
Patent No. 6448043  
GENERAL INFORMATION:  
APPLICANT: Gil H. Choi  
TITLE OF INVENTION: Enterococcus faecalis polynucleotides and polypeptides  
NUMBER OF SEQUENCES: 496  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: Human Genome Sciences, Inc.  
STREET: 9410 Key West Avenue  
CITY: Rockville  
STATE: Maryland  
COUNTRY: USA  
ZIP: 20850  
COMPUTER READABLE FORM:  
MEDIUM TYPE: Diskette, 3.50 inch, 1.44mb storage  
COMPUTER: HP Vectra 486/33  
OPERATING SYSTEM: MSDOS version 6.2  
SOFTWARE: ASCII Text  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/09/071, 035  
FILING DATE:  
CLASSIFICATION:  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER:  
FILING DATE:  
ATTORNEY/AGENT INFORMATION:  
NAME: A. Anders Brookes  
REGISTRATION NUMBER: 36,373  
REFERENCE/DOCKET NUMBER: PB369P2  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (301) 309-8504

TELEFAX: (301) 309-8512  
INFORMATION FOR SEQ ID NO: 358:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 1074 amino acids  
TYPE: amino acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
MOLECULE TYPE: protein  
US-09-071-035-358

Query Match 5.1%; Score 88.5; DB 4; Length 1074;  
Best Local Similarity 19.3%; Pred. No. 12;  
Matches 53; Conservative 40; Mismatches 87; Indels 95; Gaps 12;

QY 50 SKHFWPEV-PKXAYDMEHTFY-----SNGKKKIYMEIDVTRTEIFRSG----- 93  
DB 555 NEYFWOETKAPESYTLDETKYPSIKVDNNEKNVITR-DYTAKEQVIRFGDFPKFAG 613  
QY 94 -----NGTDETLFVHD-----FKNGYTGIFYV- 115  
DB 614 SADGTATGFNDLSEKVSPLSGTIXEITGADKATTACNEQLGPDGKFNLEPGDYLL 673  
QY 116 -----GLQKCFIKTQIKVIPSEPEEIDENEETITTFEQ-----SVIWPAPKPIE 164  
DB 674 EIEAPBGFQKI--TPLEIRSTFKENKDYAKSEVYFTITEBQKQPIKMTVPYK-- 727  
QY 165 NRDFLXSKILTEICDVTWYINPTLISVSELQ-----DFEEEGDLHFPPANE 212  
DB 728 ----LNNPFSVSLNRLMYDLPEKEDSLSLATWMDGKNKNTLDFTLVDLRLNLS 783  
QY 213 KKGIEQNEQW--VVPQYKVEKTRHARQASEELPI 245  
DB 784 IK-----EDWYVVAQAIDVEATKAAQEKDEKAPV 813

Search completed: March 30, 2004, 14:33:05  
Job time : 23 secs



GenCore version 5.1.6  
Copyright (c) 1993 - 2004 Compugen Ltd.

OM protein - protein search, using sw model

Run on: March 30, 2004, 14:29:47 ; Search time 20 Seconds  
(without alignments)  
1524.636 Million cell updates/sec

Title: US-10-089-600-2

Perfect score: 1745  
Sequence: 1 MAKPPENCDCCHILNAEAF.....VICRVIMPCNWWARMLGRV 317

Scoring table:  
BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 283366 seqs, 96191526 residues

Total number of hits satisfying chosen parameters: 283366

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database :  
1: pir1:\*  
2: pir2:\*  
3: pir3:\*  
4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	1745	100.0	317	2 JC7597	chondromodulin-1
2	1694	97.1	317	2 JC7603	tenomodulin - mouse
3	514.5	29.5	335	2 JTO569	chondromodulin-1 p
4	492.5	28.2	333	2 JTO287	chondromodulin-1 p
5	112.5	6.4	499	2 E36969	histidine kinase (
6	100.5	5.8	350	2 T34002	hypothetical prote
7	100	5.7	827	2 A90544	hypothetical prote
8	100	5.6	844	2 P86231	hypothetical prote
9	97.5	5.6	374	2 A59233	protease inhibitor
10	97.5	5.6	799	2 C82929	ATP synthase alpha
11	97	5.6	655	2 A57681	hypothetical prote
12	96.5	5.5	732	2 S73089	glycosyltransferas
13	96	5.5	2109	1 A64309	genome polypeptid
14	95.5	5.5	622	2 A64494	hypothetical prote
15	95.5	5.5	752	2 H86770	hypothetical prote
16	95.5	5.5	837	2 JNO292	antigen 332 - mala
17	95.5	5.5	1090	2 H86806	hypothetical prote
18	95	5.4	1045	1 SYEXT	isolectine-BMA 11
19	94.5	5.4	658	2 A64584	hypothetical prote
20	93.5	5.4	313	2 T19503	hypothetical prote
21	93	5.3	325	2 T18283	hypothetical prote
22	93	5.3	1429	2 T19422	hypothetical prote
23	92.5	5.3	198	2 A70013	sulfite oxidase ho
24	92	5.3	608	2 D72306	hypothetical prote
25	92	5.3	891	2 B64614	hypothetical prote
26	92	5.3	916	2 F71962	hypothetical prote
27	91.5	5.2	869	2 S76720	hypothetical prote
28	91.5	5.2	1056	1 GNLJG3	HIV-1 retropepsin
29	91	5.2	351	2 T43791	cystathionine beta

30	91	5.2	354	1 VWPJL	coat protein VP1 -
31	90.5	5.2	985	2 T41135	hypothetical prote
32	90	5.2	452	2 D64583	hypothetical prote
33	90	5.2	505	2 B64560	poly E-rich protei
34	90	5.2	595	2 E69470	amino-acid ABC tra
35	90	5.2	692	1 P3XRPC	inner layer protei
36	90	5.2	920	2 T08853	protein phosphatas
37	90	5.2	1471	2 T19506	hypothetical prote
38	89.5	5.1	1054	1 GNLJG5	HIV-1 retropepsin
39	89.5	5.1	1233	2 S64271	hypothetical prote
40	89.5	5.1	1537	2 JCA172	DNA (cytosine-5-)-
41	89.5	5.1	1965	2 T33216	hypothetical prote
42	89	5.1	431	2 I51189	IFAPA-400 - chicke
43	89	5.1	1606	2 T34073	paranem - chicke
44	89	5.1	1840	2 T25091	transitin - chicke
45	88.5	5.1	440	2 E71625	variant-specific S

## ALIGNMENTS

## RESULT 1

JC7597  
chondromodulin-1 like protein, CHML - human  
C/Species: Homo sapiens (man)  
C/Date: 30-Jun-2001 #sequence\_revision 30-Jun-2001 #text\_change 30-Jun-2001  
C/Accession: JC7597  
R/Yamana, K., Wada, H., Takahashi, Y., Sato, H., Kashihara, Y., Kiyoki, M.  
Biochem. Biophys. Res. Commun. 280, 1101-1106, 2001  
A/Title: Molecular cloning and characterization of CHML, a novel member molecule siml.  
A/Reference number: JC7597; WUID:21092728; PMID:11162640  
A/Accession: JC7597  
A/Molecule type: mRNA  
A/Residue: 1-317 <YAM>  
C/Comment: This protein is a novel type II transmembrane protein which is similar to ch  
C/Genetics:  
A/Gene: chml1  
A/Map position: Xq22  
C/Keywords: transmembrane protein  
F/31-50/Domain: transmembrane #status predicted <TMM>  
F/265-317/Region: highly conserved

Query Match 100.0%; Score 1745; DB 2; Length 317;  
Best Local Similarity 100.0%; Pred. No. 1.8e-126;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY	1	MAKNPPENCDCCHILNAEAFSKKIKSLKICGLVFGILATLIVLFMSKHPFVPPK 60			
DB	1	MAKNPPENCDCCHILNAEAFSKKIKSLKICGLVFGILATLIVLFMSKHPFVPPK 60			
QY	61	AYMEHTFYSNGEKKKIYWEIDPVTRTEIFRSNGTDETFLEVHDFKNGTGTGIFVGLQKC 120			
DB	61	AYMEHTFYSNGEKKKIYWEIDPVTRTEIFRSNGTDETFLEVHDFKNGTGTGIFVGLQKC 120			
QY	121	FITQIKVPEFSEPEEIDENEITTFEFGSVIVWPAKPIENDDFKSKILICCN 180			
DB	121	FITQIKVPEFSEPEEIDENEITTFEFGSVIVWPAKPIENDDFKSKILICCN 180			
QY	181	VIMWINPTLISVSELDQFESEGEDLHPANEEKGIQNEQWVVPQVYKTRHARQSE 240			
DB	181	VIMWINPTLISVSELDQFESEGEDLHPANEEKGIQNEQWVVPQVYKTRHARQSE 240			
QY	241	EEIPINDYENGIEPPMDDERGYCCICRGRNRYRRCCEPLIGYVPYCYQGGRVVC 300			
DB	241	EEIPINDYENGIEPPMDDERGYCCICRGRNRYRRCCEPLIGYVPYCYQGGRVVC 300			
QY	301	RVIMPCNWWARMLGRV 317			
DB	301	RVIMPCNWWARMLGRV 317			

## RESULT 2

JC7603

tenomodulin - mouse  
 C/Species: Mus musculus (house mouse)  
 C/Date: 30-Jun-2001 #sequence\_revision 30-Jun-2001 #text\_change 30-Jun-2001  
 C/Accession: J07603  
 R/Shukunani, C.; Oshima, Y.; Hiraki, Y.  
 Biochem. Biophys. Res. Commun. 280, 1323-1327, 2001  
 A/Title: Molecular cloning of tenomodulin, a novel chondromodulin-I related gene.  
 A/Reference number: J07603; MUID:21092761; PMID:11162673  
 A/Contents: Embryo, 17-days  
 A/Accession: J07603  
 A/Molecule type: mRNA  
 A/Residues: 1-317 <SHU>  
 A/Reference number: J07603  
 A/Cross-references: GB:A0219993  
 C/Comment: Tenomodulin is a type II transmembrane protein on cell surface with an antiar  
 C/Genetics:  
 C/Keywords: transmembrane protein  
 F/31-50/Domain: transmembrane #status predicted <TM>  
 F/255-317/Region: conserved carboxyl-terminus, homologous to chondromodulin-I #status pr  
 F/255-306/Domain: antiangiogenic #status predicted <ANT>

Query Match 97.1%; Score 1694; DB 2; Length 317;  
 Best Local Similarity 96.2%; Pred. No. 1.5e-122;  
 Matches 305; Conservative 6; Mismatches 6; Indels 0; Gaps 0;

QY 1 MAKPPENCDCCHILNAEAFKSKICKSLKICGLVFGIILATLIVLFWGSKHFMPEVPPK 60  
 DB 1 MAKPPENCDCCHILNAEAFKSKICKSLKICGLVFGIILATLIVLFWGSKHFMPEVSK 60  
 QY 61 AYDMEHTFVSGEKKKIYMEIDPVTETETFRSGNGDETLVHDPNKGTGIVFGLQKC 120  
 DB 61 TYDMEHTFVSGEKKKIYMEIDPVTETETFRSGNGDETLVHDPNKGTGIVFGLQKC 120  
 QY 121 FITQIKVPESEPEEIDENEITTFPEQSVIVPAEKPLENDPLKSKILIEICDN 180  
 DB 121 FITQIKVPESEPEEIDENEITTFPEQSVIVPAEKPLENDPLKSKILIEICDN 180  
 QY 181 VTWYINPTLISVSELODFEEBEGDLHFPANKEKGIQONQWVVPQVKEKTRHARQASE 240  
 DB 181 VTWYINPTLISVSELODFEEBEGDLHFPANKEKGIQONQWVVPQVKEKTRHARQASE 240  
 QY 241 EELPINDYTENGIEFPMDLDERGYCITCRGRNRCRRCCEPLGYPPYCYGGGRVTC 300  
 DB 241 EELPINDYTENGIEFPMDLDERGYCITCRGRNRCRRCCEPLGYPPYCYGGGRVTC 300  
 QY 301 RVIMPQNMWVARMIGRV 317  
 DB 301 RVIMPQNMWVARMIGRV 317

### RESULT 3

J070569  
 Chondromodulin-I precursor - bovine  
 N/Alternate names: 18k glycoprotein, cartilage; SCGP; small cartilage-derived glycoprote  
 C/Species: Bos primigenius taurus (cattle)  
 C/Date: 17-Apr-1993 #sequence\_revision 17-Apr-1993 #text\_change 05-Nov-1999  
 C/Accession: J070569; A33138; A36431  
 R/Hiraki, Y.; Tanaka, H.; Inoue, H.; Kondo, J.; Kamimono, A.; Suzuki, F.  
 Biochem. Biophys. Res. Commun. 175, 971-977, 1991  
 A/Title: Molecular cloning of a new class of cartilage-specific matrix, chondromodulin-I  
 A/Reference number: J070569; MUID:21222210; PMID:1709014  
 A/Accession: J070569  
 A/Molecule type: mRNA  
 A/Residues: 1-335 <HR>  
 A/Cross-references: GB:W65081; NID:9162840; PIRN:AAA30445.1; PID:9162841  
 A/Note: part of this sequence, including the amino end of the mature protein, was determ  
 R/Name, P.J.; Treep, J.T.; Young, C.N.  
 submitted to the Protein Sequence Database, October 1990  
 A/Reference number: A33138  
 A/Accession: A33138  
 A/Molecule type: protein  
 A/Residues: 215-220, 'W', '222', 'NE', '225-335 <NEA>  
 A/Note: 221-Val was also found

R/Name, P.J.; Treep, J.T.; Young, C.N.  
 J. Biol. Chem. 265, 9628-9633, 1990  
 A/Title: An 18-kDa glycoprotein from bovine nasal cartilage. Isolation and primary stru  
 A/Reference number: A36431; MUID:50277643; PMID:2351561  
 A/Accession: A36431  
 A/Molecule type: protein  
 A/Residues: 215-220, 'MNE', '224-225, '227-256, 'D', '257-335 <NE2>  
 A/Note: 221-Val was also found  
 A/Note: this sequence has been revised in reference A36535  
 R/Name, P.J.; Treep, J.T.; Young, C.N.  
 J. Biol. Chem. 265, 22056, 1990  
 A/Reference number: A36535  
 A/Contents: annotation; sequence revision  
 C/Comment: This protein stimulates DNA synthesis of cultured growth-plate chondrocytes  
 C/Keywords: extracellular matrix; glycoprotein; transmembrane protein  
 F/43-59/Domain: transmembrane #status predicted <TM>  
 F/215-335/Product: chondromodulin-I #status experimental <MAT>  
 F/223-236/Binding site: carbohydrate (Thr) (covalent) #status experimental  
 F/244/Binding site: carbohydrate (Asn) (covalent) #status experimental  
 F/283-287, 284-324/Disulfide bonds: (or 283-324, 284-287) #status experimental  
 F/294-318, 298-314/Disulfide bonds: #status experimental

Query Match 29.5%; Score 514.5; DB 2; Length 335;  
 Best Local Similarity 33.0%; Pred. No. 5e-32;  
 Matches 115; Conservative 60; Mismatches 109; Indels 65; Gaps 8;

QY 6 PENCEDCH--ILNFAKSKICKSLKICGLVFGIILATLIVLFWGSKHFMPEVPPKAYD 63  
 DB 15 PDDVFCSPPAYAAVYTKPSPARLKNAGAVVLSGVLLIAGIGAFYWKSGDNHITN 74  
 QY 64 MEHTFVSGEKKKIYMEIDPVTETETFRSGNGDETLVHDPNKGTGIVFGLQKCFIK 123  
 DB 75 VHYTMSINGKLQDSMEIDANNLETFMGSGAEVAVNDPFGITGIFAGGEKCYIK 134  
 QY 124 TQIKV-PEF-----SEPEEIDENEITTFPEQSVIVPAEKPLENDPLKSKILIEICDN 173  
 DB 135 AQVARIPEVGTMTKQSISSLEBGT---MPYKIEENLIVAGQPVKDNSFL-SSK 188  
 QY 174 ILEICDVTWYINPTLISVSELODFEEBEGDLHFPANKEKGIQONQWVVPQVKEKTR 233  
 DB 189 VLEIGDLPFWLKPT-----YF---KEIQEREELVRKIVTTT 226  
 QY 234 HARQASEELPINDYTENG-----IEEDPMLDERGYCIT 266  
 DB 227 RLRSFGPGTPAPGRPNNGTRPSVQEDAEPPNPDPYHQBESMTDPDLDEHICIE 286  
 QY 269 CRGRNRCRRCCEPLGYPPYCYGGGRVLCRYIMPQNMWVARMIGRV 317  
 DB 287 CRGSYTHQKICEPLGIVHPMPYNYGCRSACRTYIMCSMWVARIIGMV 335

### RESULT 4

J020287  
 Chondromodulin-I precursor [similarity] - rat  
 C/Species: Rattus norvegicus (Norway rat)  
 C/Date: 20-Apr-2000 #sequence\_revision 20-Apr-2000 #text\_change 20-Apr-2000  
 C/Accession: J020287  
 R/Shukunani, C.; Hiraki, Y.  
 Biochem. Biophys. Res. Commun. 249, 865-890, 1998  
 A/Title: Expression of cartilage-specific functional matrix chondromodulin-I mRNA in ra  
 A/Reference number: J020287; MUID:98401184; PMID:9731231  
 A/Accession: J020287  
 A/Molecule type: mRNA  
 A/Residues: 1-333 <SHU>  
 A/Cross-references: DDBJ:AF072129; NID:93641531; PIRN:AAC36440.1; PID:93641532  
 C/Keywords: extracellular matrix; glycoprotein; transmembrane protein  
 F/43-59/Domain: transmembrane #status predicted <TM>  
 F/215-333/Product: chondromodulin-I #status predicted <MAT>  
 F/223/Binding site: carbohydrate (Thr) (covalent) #status predicted  
 F/243/Binding site: carbohydrate (Asn) (covalent) #status predicted  
 F/261-265, 262-322, 292-316, 296-312/Disulfide bonds: #status predicted

Query Match 28.2%; Score 492.5; DB 2; Length 333;

Best Local Similarity 33.4%; Pred. No. 2,4e-30;  
Matches 110; Conservative 72; Mismatches 120; Indels 27; Gaps 12;

```

QY 6 PENCEDCHIAAFAFSKTI-----CKSLKICGLVFGIILALIVLFW--GSKHFWPEVP 58
DB 15 PDVEFC--GPPAVAVIVKPSGPAFLKVGAVV--LISGALLLFGAIGAYLWKSGD 69
QY 59 KKAYDMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLLEHDFKNGYTIYFVGLQ 118
DB 70 NHIVNHYMTSINGKLQDSMEIDARNNLETFFMGSSAEALIVNFGNGITIRFAGE 129
QY 119 KCFIKTQIKY-IPBF-SEPEEID--ENBEITTFEFGSVIVPAKPIENDFLKNK 173
DB 130 KCVIKAKVAKRVPVGVITQOISISELEGKIMPKHEEALVWVAGQPVQDNSFL-SAR 188
QY 174 ILLECDVAVTWVWVWIPPLISVSELODFEEGEDLHFPANEK--GIEQNEQWVVPQVKE 230
DB 189 VLEICGDLPIFWLKPITY--KREIQRERREYVAKTIVPTTKRPSGPRGNG--PALMRD 244
QY 231 KTRHARQASEEELPINDYTENG--IEPDPMLDERGYCCICRGNRCREVECPPLIGYYP 288
DB 245 SRPESVQDSPEPNDNFYHOGESMTDPRLDHEGICICRPSRYTHCQKICEPLGQYNP 304
QY 289 YPYCYGGRVYICRYMPCMMVWARMGRV 317
DB 305 WPIYVQGRSACRYVWPCSMWVARIIGMV 333

```

## RESULT 5

histidine kinase (HAMP, Hiskra and HATPase domains) (imported) - Clostridium acetobutylicum  
C:Species: Clostridium acetobutylicum  
C:Date: 14-Sep-2001 #sequence\_revision 14-Sep-2001 #text\_change 14-Sep-2001  
R:Noiling, J.; Breton, G.; Omeichenko, M.V.; Markarova, K.S.; Zeng, Q.; Gibson, R.; Lee, J.; Daly, M.J.; Bennett, G.N.; Koonin, E.V.; Smith, D.R.  
J. Bacteriol. 183: 4823-4838, 2001  
A:Title: Genome Sequence and Comparative Analysis of the Solvent-Producing Bacterium C1C  
A:Reference number: A96900; MUID:21359325; PMID:21359325  
A:Accession: E96969  
A:Status: preliminary  
A:Molecule type: DNA  
A:Residues: 1-499 <KUR>  
A:Cross-references: GB:AE001437; PIDN:AAK78544.1; PID:G15023432; GSPDB:GN00168  
A:Experimental source: Clostridium acetobutylicum ATCC824  
C:Genetics:  
A:Gene: CAC0565

Query Match 6.4%; Score 112.5; DB 2; Length 499;  
Best Local Similarity 22.3%; Pred. No. 0.59;  
Matches 71; Conservative 49; Mismatches 103; Indels 95; Gaps 21;

```

QY 15 LNAFAFSKTKCKSLIKI-CGLVFGIILALIVL-----FMGSKHFWPEVP----- 58
DB 9 LKLYSFSKXKVKSRILELVTFGICLAFILCSWYTGYYNDGMAAEVDYSSGIGRIS 68
QY 59 -----KKAYDMEHTFYSNGEKK--KIYMEIDPVTRTE--IFRSNGTDETLLE 101
DB 69 SDMEDKLMDLQKSSDEINKMIDSGACDDIKVYL-----TDTGKYLFFSSNKEKID 123
QY 102 VHD-----FRNGYTGIVFVGLQKCFIKTIQIKVIPFSEPEEID--ENEE---ITT 147
DB 124 IHDLVKSEKFEKREYKATVINTVNGIKITISL--SEVEQYNIDGIKIDNKAVYIVS 181
QY 148 TFFEGSVIVW-PAKPIENRDFLKNKSLLECDNVWVW-----INPLTISVS 194
DB 182 GIPKOKTIVIRPGVSP---FL--SGIIAVITTFVFIPLTNKMEYBSVSGVLEIS 234
QY 195 ELQ--DFE--EEGEDLHFPANEKKGIEQNEQWVP--QVKEKTRHARQASEEELPINDYT 249
DB 235 KGNLDYRILMLMGED-----ELKNLADNINSMASELQRIKERBAEKTK-----NDLI 282
QY 250 ENGIEFD---PMLDERGY 264

```

DB 283 TN-VSHDIRPLETSYKGY 299

## RESULT 6

hypothetical protein W03G1.2 - Caenorhabditis elegans  
C:Species: Caenorhabditis elegans  
C:Date: 29-Oct-1999 #sequence\_revision 29-Oct-1999 #text\_change 29-Oct-1999  
C:Accession: T34002  
R:Pauley, A.; Scheet, P.; Harper, M.  
submitted to the EMBL Data Library, February 1999  
A:Description: The sequence of C. elegans coemid W03G1.  
A:Reference number: Z21454  
A:Accession: T34002  
A:Status: preliminary; translated from GB/EMBL/DDBJ  
A:Molecule type: DNA  
A:Residues: 1-350 <PAU>  
A:Cross-references: EMBL:AF125964; PIDN:AAID14750.1; GSPDB:GN00022; CESP:W03G1.2  
A:Experimental source: strain Bristol N2; clone W03G1  
C:Genetics:  
A:Gene: CESP:W03G1.2  
A:Map position: 4  
A:Introns: 19/3; 97/1; 233/3; 335/3

Query Match 5.8%; Score 100.5; DB 2; Length 350;  
Best Local Similarity 20.9%; Pred. No. 3.1;  
Matches 52; Conservative 40; Mismatches 72; Indels 85; Gaps 13;

```

QY 50 SGHFWPEVPRKAY-----DMEHTFY-SNGEKKIYMEIDPVTRTEIFRSNGT 96
DB 145 SKLMDRVPKPKYRLKNKPDGMMLCDEPSTFVRTTSKKKXLSLPVE-----DSY 196
QY 97 DETLEVHDFRNGYTGIVFVGLQKCFIKTIQIKVIPFSEPEEIDNEEITTFEFGSVIV 156
DB 197 DVMRKLEEDK-----KEPEQVYRGQDVPF-----W 222
QY 157 VBAEKPIENRDFLKNKSLIKI-CDNVTWY-----WINPLISVSELODFEE--EGED 205
DB 223 AKRIETTD-DKMEIDEATVSTDLHWRNKKLSLPIKLVLNELQPLVELKKDE 281
QY 206 LHPANEKKGIEQNEQWVVPQVKEKT---RHARQASEEELPINDYTENGIEFPMLDE 261
DB 282 VHF-----ETGLVFSNTIRSLILVHESARALSDGYD--DEYSDKVVQFDTQEP 329
QY 262 RGYCCICYR 270
DB 330 ATY--TYSR 336

```

## RESULT 7

hypothetical protein WYPU 2570 (imported) - Mycoplasma pulmonis (strain UAB CTIP)  
A90544  
C:Species: Mycoplasma pulmonis  
C:Date: 24-May-2001 #sequence\_revision 24-May-2001 #text\_change 03-Aug-2001  
C:Accession: A90544  
R:Chamand, I.; Helling, R.; Ferris, S.; Barbe, V.; Samson, D.; Galisson, F.; Moszer, I.; Nucleic Acids Res. 29, 2145-2153, 2001  
A:Title: The complete genome sequence of the murine respiratory pathogen Mycoplasma pul  
A:Reference number: A99512; MUID:21267165; PMID:11353084  
A:Accession: A90544  
A:Status: preliminary  
A:Molecule type: DNA  
A:Residues: 1-827 <KUR>  
A:Cross-references: GB:AL445566; PID:G14089670; PIDN:CAC13430.1; GSPDB:GN00153  
A:Experimental source: strain UAB CTIP  
C:Genetics:  
A:Gene: WYPU 2570  
A:Genetic code: SGC3  
C:Superfamily: valine-tRNA ligase

Query Match 5.7%; Score 100; DB 2; Length 827;  
Best Local Similarity 21.8%; Pred. No. 10;

Matches 57; Conservative 45; Mismatches 67; Indels 92; Gaps 14;

QY 24 KICKSLKIGLVGIALTLIVLFWSKFWPEVP-----KKAYD---MHTYVSGE 73  
 DB 634 KITKNTDPQKIVFKELLIL-----HPIPLTTHLTKIYDGLIHHTF---E 679  
 QY 74 KKKIYWEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLOKCFIKTOIKVPEPS 133  
 DB 680 NKKL-----KTQ-----KNVVDQLIV-----IRAIRSEFR 704  
 QY 134 EPEEIDEENBITTTTFEQSVIWPV---AEKPIENRDL-----KNSKILEICDNTM 183  
 DB 705 E-KYQSKKEKIK-----YWIQCQFLQEDIDAINFLTSELSQNSMTIVENIKI 755  
 QY 184 YWNPFLISVSELODEEGEDLHPANEKGIQONQWV--VPQVKEKTRHARQASEE 241  
 DB 756 FMILPKNIEENLSKEKAKKEFLKFEIKRAQSLLNEKRTISKAPFLKVE-----E 806  
 QY 242 ELPINDYENGIFPMDLDER 262  
 DB 807 KATLEKYQ--LQKLLEDEK 824

## RESULT 8

F86231 hypothetical protein [imported] - Arabidopsis thaliana

C/Species: Arabidopsis thaliana (mouse-ear cress)

C/Date: 02-Mar-2001 #sequence\_revision 02-Mar-2001 #text\_change 31-Mar-2001

C/Accession: F86231

R/Proteolysis: A.; Ecker, J.R.; Palm, C.U.; Federspiel, N.A.; Kaul, S.; White, O.; Alonso, Chn, C.W.; Chung, M.K.; Conn, L.; Conway, A.B.; Conway, A.R.; Creasy, T.H.; Dewar, K.; Jensen, N.F.; Hughes, B.; Hultzer, L.

Nature 408, 816-820, 2000

A/Authors: Hunter, J.L.; Jenkins, J.; Johnson-Hopson, C.; Khan, S.; Khaykin, E.; Kim, C.C.A.; Li, J.H.; Li, Y.; Liu, X.; Liu, S.X.; Liu, Z.A.; Luros, J.S.; Maiti, R.; Marziani, Rizzo, M.; Rooney, T.; Rowley, D.; Sakano, H.

A/Authors: Salzberg, S.L.; Schwartz, J.R.; Shinn, P.; Southwick, A.M.; Sun, H.; Tallon, Ker, M.; Wu, D.; Yu, G.; Fraser, C.M.; Venter, J.C.; Davis, R.W.

A/Title: Sequence and analysis of chromosome 1 of the plant Arabidopsis.

A/Reference number: A86141; MIMD:21016719; PMID:11130712

A/Accession: F86231

A/Status: Preliminary

A/Molecule type: DNA

A/Residues: 1-844 <STO>

A/Cross-references: GB:AE005172; NID:g2160167; PIDN:AA860730.1; GSPDB:GN00141

C/Genetics:

A/Map position: 1

C/Superfamily: myb-related protein cdc5; myb DNA-binding repeat homology

Query Match 5.7%; Score 100; DB 2; Length 844;

Best Local Similarity 22.7%; Pred. No. 10;

Matches 58; Conservative 36; Mismatches 88; Indels 74; Gaps 12;

QY 90 FREGNTDTELEHND-----FKNGYGIYFVGLQCKFIQTQIKVLP---E 131  
 DB 447 FRELHINDMDHESAKLERQREBARSLSGLT---GLPQKNEYQIVAPPPPE 501  
 QY 132 FSEPEEIDENBEITTFEQSVIWPVPAKPIENRDL--KNSKILE----- 176  
 DB 502 SEPEPEKIEED-----MSDRARAKAEBAKQGLLKKRKYVLRDLPKPPASLAVI 554  
 QY 177 ----ICDNTWYINPTLISVSELODFEE---EGEDLHPANEKGIQONQWVVPQ 226  
 DB 555 RNSILSADGDKSSVPTPEVADKRVRELLQLLEHDAKAPLDDK-----AEK 604  
 QY 227 VKVETRHRARQASEEELPINDYTENGI--EEDPMLDERG--YCCYICRGRNR-----YGR 278  
 DB 605 KKKAKKRTKRSAS-QVLAIDDFDENELQADKKIKKKEGFLCVSMGHEKTLDDVVEAHN 663  
 QY 279 VCEPLIGYPPYCYQ 294  
 DB 664 TCVNDIMYFPTRSAE 679

## RESULT 9

A59273 proteinase inhibitor 8 - human

N/Alternate names: cytoplasmic antiproteinase 2 (CAP-2)

C/Species: Homo sapiens (man)

C/Date: 02-Jun-2000 #sequence\_revision 02-Jun-2000 #text\_change 15-Sep-2003

C/Accession: A59273

R/Sprecher, C.A.; Morgenstern, K.A.; Mathewes, S.; Dahlen, J.R.; Schrader, S.K.; Foster

J. Biol. Chem. 270, 29854-29861, 1995

A/Title: Molecular cloning, expression, and partial characterization of two novel membe

A/Reference number: A59273; MIMD:96102039; PMID:8530382

A/Accession: A59273

A/Status: Preliminary; not compared with conceptual translation

A/Molecule type: mRNA

A/Residues: 1-374 <SPR>

A/Cross-references: GB:L40377; NID:g1160926; PIDN:AA41939.1; PID:g1160927

A/Experimental source: tissue type placenta; note (vector lambda gtl1); gene CAP2

C/Genetics:

A/Gene: GDB:P18; CAP2

A/Cross-references: GDB:599392; OMIM:601697

A/Map position: 18q21.3-18q21.3

C/Superfamily: Serpin

C/Keywords: proteinase inhibitor

F.339/Inhibitory site: Arg (undentified proteinase) #status predicted

Query Match 5.6%; Score 97.5; DB 2; Length 374;

Best Local Similarity 21.3%; Pred. No. 5.8;

Matches 63; Conservative 42; Mismatches 112; Indels 79; Gaps 14;

QY 7 ENCEDC--HILNAPFKSKICKSLKICGVFGIALTLI--VLFWSKHFVPEPKAY 62  
 DB 123 EDTECKKHINDVAKTEBKISVDAGVDPITKLVNAYLFCK--VNEQFPKRY 179  
 QY 63 DMEHTFYSNGEKKIYWEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLOKCFI 122  
 DB 180 TRGMLFKTNEKKIKVQK-----MFEKAK-----FPMGYAD-----EV 211  
 QY 123 KTOIKVLPSEPEPEE-----DENEETTT-----FEGSVIWPVPAKPIENR--DFL 169  
 DB 212 HTQVLELPYV---EELSMTILPDDNTDULAVVEKALTVEKFAWTMSKLTYSKQVFL 268  
 QY 170 KNSKILEICDNTWYINPTLISVSELODEEGEDLHPANEK-----GIEON 219  
 DB 269 PRLKLE-----SYDLEPLRLGIMIDAEKADPSGKSTKQVPLSKVAHKCFEYEVN 322  
 QY 220 EQWVVPQVKEKTRHARQASEEELPINDYTENGEIPEPML-----DERGYCICRR 271  
 DB 323 EEGTEAATAATVAVNSKRSNRPFCADH-----FLPFTIRRHKTNCILFCGR 370

## RESULT 10

C82929 aryl synthase alpha chain U0132 [imported] - Ureaplasma urealyticum

C/Species: Ureaplasma urealyticum

C/Date: 18-Aug-2000 #sequence\_revision 20-Aug-2000 #text\_change 20-Aug-2000

C/Accession: C82929

R/Glass, J.I.; Letkowitz, E.J.; Glass, J.S.; Hainer, C.R.; Chen, E.Y.; Cassell, G.H.

submitted to Genbank, February 2000

A/Description: The complete sequence of Ureaplasma urealyticum: Alternate views of a mi

A/Reference number: A82670

A/Accession: C82929

A/Status: Preliminary

A/Molecule type: DNA

A/Residues: 1-799 <GLA>

A/Cross-references: GB:AE002114; GB:AF222894; NID:g6899086; PIDN:AAAF30538.1; GSPDB:GN00:

A/Experimental source: serovar 3; biovar 1

C/Genetics:

A/Gene: atpA-2; U0132

A/Genetic code: SGC3

Query Match 5.6%; Score 97.5; DB 2; Length 799;

Best Local Similarity 20.7%; Pred. No. 15;

Matches 57; Conservative 42; Mismatches 79; Indels 97; Gaps 12;

QY 44 IYFMGSKHFWPVP-----KKAYMEHT----- 67  
 Db 436 IITFTYKKEKIPVPERIDDFEYLLNYFKGTKLADLEDKAPKENTPAFRCALQCA 495  
 QY 68 ---FYSNGEKKKIYMEIDPVTRTEIFRSGNGTDETELVHDFKNGYGIYVGLQKCFIKT 124  
 Db 496 INSFLNNSODFKPCEAEQOTAPDKPFVE---NESIVV-DGENDFN-----FINE 540  
 QY 125 QIKVIP-----EFSEPEEIDENEITTFEEQSVIWPAPKPI----- 163  
 Db 541 EVSIKPTTTESEAVQIEKQDVEFEPQELITNKMEBNHFEF---VEPEKLTICENHEF 596  
 QY 164 ---ENRDFLNKSKILEICDNVWYMINPTLISVSELODFEEBEGDLH--FPANEEKGIE- 217  
 Db 597 EIENQSKIEGQVLE--DINHEYSIYETVEQSGEVDNDSKODDLEVLVPVIEIHDBA 654  
 QY 218 ----QNEQVY----VPQVKEKTRHARQASEE 242  
 Db 655 ILDERENRNVVFSDSAVSEVKOTIMISSNEAE 689

RESULT 11  
 A57681  
 hypothetical protein - Mycoplasma capricolum  
 C/Species: Mycoplasma capricolum  
 C/Date: 26-Jul-1996 #sequence\_revision 26-Jul-1996 #text\_change 07-Dec-1999  
 C/Accession: A57681  
 R/Zhu, P.P.; Reiter, J.; Peterkofsky, A.  
 Protein Sci. 3, 2115-2128, 1994  
 A/Title: Unique dicistronic operon (psti-crr) in Mycoplasma capricolum encoding enzyme I  
 ncing, promoter analysis, and protein characterization.  
 A/Reference number: A57681; MUID:95218642; PMID:7703858  
 A/Accession: A57681  
 A/Status: preliminary; translated from GB/EMBL/DBJ  
 A/Molecule type: DNA  
 A/Residues: 1-655 <RES>  
 A/Cross-references: EMBL:U5110; NID:g602679; PID:AAA70404.1; PID:g602684  
 C/Genetic code: SGC3

Query Match 5.6%; Score 97; DB 2; Length 655;  
 Best Local Similarity 21.3%; Pred. No. 13;  
 Matches 57; Conservative 44; Mismatches 99; Indels 68; Gaps 12;

QY 7 ENEDCHILNAEAFKSKICKSLKICGVGIIALLILYIFMSKPFVPEVKKAYDMEH 66  
 Db 155 DNEEDIDDSKLEILSKYI-----DNQHNYPDYKRN----- 185  
 QY 67 TFSNGEKKKIYMEIDPVTRT-----EIFRSGNGTDETELVHDFKNGYGIYVGL 116  
 Db 186 --FKTVAEIEYKELVYRTSIFKIVYKDGGLSNOTGWLDDYHKXSN--TNKY---- 238  
 QY 117 LQKCFITQKIVPESEPEEIDEENETTFEEQSV--VIVPAEKPIENRDLPLANSK 173  
 Db 239 --KMFATINLHVLADEFNSLTLD--EQNKEF--NYVDSGNKVIYLGKADNVTDFSRKNN 293  
 QY 174 ILIEICDNVWYMINPTLISVSELODFEEBEGDLHFPANE--KKGIEQNEQVVPVQVVEKT 232  
 Db 294 NKSSENNIANYILNN-----QDFENYKNDPFWVNSKSKGISR--PKIVGADVPMKD 344  
 QY 233 R---HARQASEEELPINDYTENGIEPD 256  
 Db 345 RAIKNHYEALQKEAINYYNKKNNNEIN 372

RESULT 12  
 S73089  
 glycosyltransferase (EC 2.4.1.-) - Sulfolobus solfataricus  
 N/Alternate names: protein c0622  
 C/Species: Sulfolobus solfataricus  
 C/Date: 09-Oct-1997 #sequence\_revision 24-Oct-1997 #text\_change 17-Mar-1999

C/Accession: S73089  
 R/Sensen, C.W.; Klenk, H.P.; Singh, R.K.; Allard, G.; Chan, C.C.Y.; Liu, Q.Y.; Penny, S.  
 Mol. Microbiol. 22, 175-191, 1996  
 A/Title: Organizational characteristics and information content of an archaeal genome: 1  
 A/Reference number: S73076; MUID:97055432; PMID:8899719  
 A/Accession: S73089  
 A/Status: nucleic acid sequence not shown; translation not shown  
 A/Molecule type: DNA  
 A/Residues: 1-732 <SEN>  
 A/Cross-references: EMBL:Y08256; NID:g1707679; PID:e284015; PID:g1707701  
 A/Experimental source: strain P2  
 A/Note: the nucleotide sequence was submitted to the EMBL Data Library, September 1996  
 C/Keywords: glycosyltransferase; hexosyltransferase

Query Match 5.5%; Score 96.5; DB 2; Length 732;  
 Best Local Similarity 20.4%; Pred. No. 16;  
 Matches 64; Conservative 51; Mismatches 114; Indels 85; Gaps 15;

QY 37 GILMTLIVLWGSNHFVPPKAYDMHTFYSN-----GEKKIYMEIDPVTRT 87  
 Db 393 GIMRL-----QQYPAIFAKGYEDTLFIYNRILSLNEVSGDLRRLSIDDFNHF 443  
 QY 88 EIFRSGNGTDETELVHDFKNGYGIYVGLQKCFIKQIKVIFPSEPEE----- 138  
 Db 444 NQSRNVTISMNTLSTHDTKFSED-----VRAISVLSIEPKMEERVIYHDDL 492  
 QY 139 ---IDENEITTFEEQSVIWPAPKPIENRDLPLANSKILEICD--NVTMYMINPTLISVS 194  
 Db 493 RPNIDKNDYR--FYQTL--VGSYEGFDNKRKIKHMKIVIRAKVHTTENPIEYEN 547  
 QY 195 ELQDFEEEG-EDLHPANEEKGIEQNEQV-----VPQV--KYEKTRH 234  
 Db 548 KYLDIDDAFENSNN--RDPFSFEKVIYFPYMSVATTLKFLSPGVPDIYQGEVWRF 606  
 QY 235 ARQASEEELPI-----NDYENGIETP--DPMLEDERGICCIYCRGRNRYCRVCEPL 284  
 Db 607 LLTDPDNMPPVDPFKKRLNLTENKLELSDPRVK-----MLVYKLLQDRREY--SLN 659  
 QY 285 GYEPYPCYOGGRV 298  
 Db 660 DYKPLPFGFORQKV 673

RESULT 13  
 A46309  
 genome polynuclein - vesicular stomatitis New Jersey virus (strain Ogden)  
 N/Alternate names: RNA nucleotidyltransferase (RNA-directed); RNA replicase  
 N/Comments: RNA-directed RNA polymerase (EC 2.7.7.48)  
 C/Species: vesicular stomatitis New Jersey virus  
 C/Date: 17-Feb-1994 #sequence\_revision 17-Feb-1994 #text\_change 11-Jun-1999  
 C/Accession: A46309  
 R/Barik, S.; Rud, E.W.; Luk, D.; Banerjee, A.K.; Kang, C.Y.  
 Virology 175, 332-337, 1990  
 A/Title: Nucleotide sequence analysis of the L gene of vesicular stomatitis virus (New J  
 A/Reference number: A46309; MUID:90177235; PMID:2155516  
 A/Accession: A46309  
 A/Molecule type: genomic RNA  
 A/Residues: 1-2109 <BAR>  
 A/Cross-references: GB:M29788; NID:g336029; PID:AAA48442.1; PID:g336030  
 C/Genetics:  
 A/Genes: L  
 C/Superfamily: Thadovirus L protein  
 C/Keywords: nucleotidyltransferase; RNA binding; RNA biosynthesis; RNA replication  
 F/530-549/Region: RNA binding #status predicted

Query Match 5.5%; Score 96; DB 1; Length 2109;  
 Best Local Similarity 22.1%; Pred. No. 67;  
 Matches 52; Conservative 46; Mismatches 91; Indels 46; Gaps 13;

QY 55 PEVPRKAYDMHTFYSNGEKKKIYMEIDPV--TRTEIFRSGNGTDETELVH--DFKN--GYTG 111  
 Db 306 PKIPKPFEEHV--KGSVQELTORSNRIOTLVDLIMSKMDVLLVLYVGSFRHWGHPF 362

QY 112 I-YFVGLQKCFIKTQIKVIPSEPEEEDEN--EETITTF-----FEQSVIYW--- 157  
 Db 363 IDYFQLEK--LHTQVNM-----EKHIDKEYPQOLASDLARLVNKKFSSSKMFPDP 413  
 QY 158 -----PAEKPIENRDFLKNKILFICNVNTMYWNPFLISVSELQDFEEEG---EDLH 207  
 Db 414 SKNSPGRPFYEVHVNKTWPTAKIOPFGDN---WKKPLICFPIPLIDPSVIYSKKS 469  
 QY 208 PANENKQIE---QNEQWVPQVKEKTRHARQASEELPINDYENGIEPDPML 259  
 Db 470 HSMNKKEVIOHRSKNKNIPIPSNKVLQITLTNRAITWKAFLDIDENGDDDDLI 524

## RESULT 14

A64494  
 Hypothetical protein MJ1554 - Methanococcus jannaschii

C/Species: Methanococcus jannaschii  
 C/Date: 13-Sep-1996 #sequence\_revision 13-Sep-1996 #text\_change 21-Jul-2000

C/Accession: A64494  
 R/Bult, C.J.; White, O.; Olsen, G.J.; Zhou, L.; Fleischmann, R.D.; Sutton, G.G.; Blake,  
 ; Reich, C.I.; Overbeek, R.; Kirkness, E.F.; Weissbrock, K.G.; Merrick, J.M.; Glodek, A.;  
 Ison, J.D.; Sadow, P.W.; Hama, M.C.; Cotton, M.D.; Roberts, K.M.; Hurst, M.A.  
 Science 273, 1058-1073, 1996

A/Authors: Kaine, B.P.; Borodovsky, M.; Klenk, H.P.; Fraser, C.M.; Smith, H.O.; Woese, C

A/Title: Complete genome sequence of the methanogenic archaeon, Methanococcus jannaschii

A/Reference number: A64300; PMID:96337999; PMID:8688087

A/Accession: A64494

A/Status: preliminary; nucleic acid sequence not shown; translation not shown

A/Molecule type: DNA

A/Residues: 1-622 <BUL>

A/Cross-references: GB:U67596; GB:L77117; NID:G2826430; PIDN:AAB99574.1; PID:G1500447; T

C/Genetics:

A/Map position: FOR:529328-1531196

C/Superfamily: hypothetical protein MJ1554

Query Match 5.5%; Score 95.5; DB 2; Length 622;  
 Best Local Similarity 22.0%; Pred. No. 16;  
 Matches 56; Conservative 40; Mismatches 79; Indels 79; Gaps 13;

QY 21 KSKTKCKSKIKIGLVGILATLILVFGSKHFW---PEVPKAYDMEHFYSNGEKKK 76  
 Db 132 KYAKIKIKNISBCGYLV-LTNNTLIVISWNTKITSYVNSNPEMKITQWD---LNGS--- 183  
 QY 77 IYMEIDPVTRTEIFRSGNGT-----DETLEVHDFKNGYTGIVF----- 114  
 Db 184 -YVD-----SRLVNGTLVYVRKNSIDCPVWNNYKIGYDKYIPELPIYSMDPD 233  
 QY 115 -----VGLQKCFIKTQIKVIPSEPEE-----EEIDENEIITTFEQSVI 155  
 Db 234 TYIISRLINIKSGKVENSTAIIVGNKTTILYMSKNLKYFAVNLKINEKLMNPLNESADK 293  
 QY 156 WVPAR-----KPIENRDFLKNKILFICNVNTMYWNPFLISVSELQDFEE 202  
 Db 294 YFPTLVADKIKRVINEEDGDNKAFVEIETIERYL--SSLPSKRNLMKKLQNDPENY 351  
 QY 203 GEDLHFPANKEKGI 216  
 Db 352 LEE-HWEPEPYTGI 364

## RESULT 15

H86770

Hypothetical protein y11b [imported] - Lactococcus lactis subsp. lactis (strain IL1403)

C/Species: Lactococcus lactis subsp. lactis

C/Date: 23-Mar-2001 #sequence\_revision 23-Mar-2001 #text\_change 03-Aug-2001

C/Accession: H86770

R/Bolotin, A.; Wincker, P.; Mauger, S.; Jallion, O.; Malarme, K.; Weissbach, J.; Earl,

Genome Res. 11, 731-753, 2001

A/Title: The complete genome sequence of the lactic acid bacterium Lactococcus lactis s

A/Reference number: AB6625; PMID:12135186; PMID:11337471

A/Accession: H86770

A/Status: preliminary

A/Molecule type: DNA  
 A/Residues: 1-752 <STO>  
 A/Cross-references: GB:AB005176; PID:G12724133; PIDN:AAK05266.1; GSPDB:GN00146  
 A/Experimental source: strain IL1403  
 C/Genetics:  
 A/Gene: y11b

Query Match 5.5%; Score 95.5; DB 2; Length 752;  
 Best Local Similarity 21.7%; Pred. No. 20;  
 Matches 63; Conservative 37; Mismatches 95; Indels 95; Gaps 17;

QY 16 NAEFKSKIKCKSKIKIGLV---EGILATLILVFGSKHFWPEVPKAYDMEH-- 66  
 Db 174 NTQILPILTWCKAKIKELCPVSKAIQUGNFITRISLIGLHFPWNYIDSMILYDGENL 233  
 QY 67 -TFYSGEKKKIYMEID-----FVTRTEIFRSQ--NGIDETLEVHDFKNGYTGIVF 114  
 Db 234 SSFLOLENNKDYSNINLVYLDHNRILKNEIVESKTINSLPDPDGFDFSKYFTDNLW 293  
 QY 115 VGL---QKCFIKTQIKVIPSEPEE-----EPEEIDENEIITTFEQSVI 157  
 Db 294 IPYTSACVKNKSTFCTIPNASGKTRHMPAKKVENMEVYKERYGISHSFVDEFTL-V 352  
 QY 158 PAEKPI-----ENRD-----FLK-----NSKILEI 177  
 Db 353 PKKQIKLILDMQKDISWYCTRFSLTLADTTQILRKGGCSNTQFLESYNQAVLDL 412  
 QY 178 CD-NTVMYWNPLISVSF-----LQDFEEGDLHFPANE 212  
 Db 413 MDKVIDAMIEPININCFEAGISVHLFPMTGPPTETLEAKNTYHF-ANE 461

Search completed: March 30, 2004, 14:33:39  
 Job time : 22 secs

GenCore version 5.1.6  
Copyright (c) 1993 - 2004 CompuGen Ltd.

# OM protein - protein search, using sw model

Run on: March 30, 2004, 14:29:47 ; Search time 17 Seconds  
(without alignments) 970.955 Million cell updates/sec

Title: US-10-089-600-2  
Perfect score: 1745  
Sequence: 1 MAGNPPENCDCIILNBEAF.....VICRVIIMPCNMTVARYMLGRV 317

Scoring table: BIOSOM62  
Gapop 10.0 , Gapext 0.5

Searched: 141681 seqs, 52070155 residues  
Total number of hits satisfying chosen parameters: 141681

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : SwissProt\_42:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	1745	100.0	317	1	TNMD_HUMAN
2	1694	97.1	317	1	TNMD_MOUSE
3	1657	95.0	317	1	TNMD_RAT
4	553	31.7	347	1	CHM1_CHICK
5	529	30.3	334	1	CHM1_RAT
6	518	29.7	334	1	CHM1_MOUSE
7	514.5	29.6	334	1	CHM1_HUMAN
8	492.5	28.2	333	1	CHM1_BOVIN
9	403.5	23.1	333	1	CHM1_RABIT
10	97.5	5.6	374	1	CHM1_BRARE
11	97	5.6	374	1	SPB8_HUMAN
12	96	5.5	655	1	YKDA_MYCA
13	96	5.5	2109	1	RRLP_VSVUO
14	95.5	5.5	622	1	YF54_MENUA
15	95	5.4	312	1	SPR2_CAREL
16	95	5.4	1044	1	SYI_MERTM
17	94.5	5.4	681	1	MP10_HUMAN
18	93	5.3	632	1	KU70_CHICK
19	93	5.3	682	1	TDR5_HUMAN
20	92.5	5.3	684	1	HTPG_PORGI
21	92	5.3	664	1	PARF-STARP
22	91.5	5.2	1056	1	POL_SIVWI
23	91	5.2	354	1	COAI_POUVC
24	90	5.2	692	1	VP3_ROTIC
25	89.5	5.1	1054	1	POL_SIVWK
26	89.5	5.1	1233	1	YF16_YEAST
27	89.5	5.1	1537	1	DNM1_CHICK
28	89	5.1	864	1	41_HUMAN
29	89	5.1	1001	1	RPGR_MOUSE
30	88	5.0	351	1	FEN_SURTO
31	87.5	5.0	149	1	RNP_MUSPA
32	87.5	5.0	634	1	YG74_MENUA
33	87	5.0	1409	1	COP1_DROME

34	87	5.0	2820	1	NPI_RAT	P97526	rattus norv
35	86.5	5.0	283	1	Y635_MENUA	Q58052	methanococ
36	86.5	5.0	699	1	H582_ARATH	P55737	arabidopsis
37	86.5	5.0	699	1	H583_ARATH	P51818	arabidopsis
38	86	4.9	606	1	SPB4_YEAST	P25808	saccharomyc
39	86	4.9	700	1	HS9C_DICDI	P54651	dictyosteli
40	85.5	4.9	493	1	ECX1_MENUA	Q80118	methanococ
41	85.5	4.9	660	1	STR_MOUSE	Q96019	mus musculu
42	85.5	4.9	661	1	STR_CRITO	P37880	cricetus
43	85	4.9	350	1	IPYR_CAREL	Q16880	caenorhabdi
44	85	4.9	414	1	Y288_MYCCE	P47530	mycoplasma
45	85	4.9	442	1	TIG_BUCAL	P57546	buchnera ap

## ALIGNMENTS

RESULT 1  
TNMD\_HUMAN STANDARD; PRT; 317 AA.  
AC Q9H2S6; Q9H2X0; Q9UTG0;  
DT 16-OCT-2001 (Rel. 40, Created)  
DT 16-OCT-2001 (Rel. 40, Last sequence update)  
DT 15-MAR-2004 (Rel. 43, Last annotation update)  
DE Tenomodulin (Tnm) (Hnm) (Chondromodulin-I like protein) (CHM1L)  
DE (hChm1L) (Myodulin).  
GN TNMD OR CHM1L.  
OS Homo sapiens (Human).  
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
OC Mammalia; Eutheria; Primates; Catarrhini; Homiidae; Homo.  
OX NCBI\_TaxID=9606;  
RN [1]  
RP SEQUENCE FROM N.A.  
RX MEDLINE=21092728; PubMed=1162640;  
RA Yamana K., Wada H., Takahashi Y., Sato H., Kasahara Y., Kiyoki M.;  
RT "Molecular cloning and characterization of Chm1L, a novel membrane  
RT molecule similar to chondromodulin-I";  
RL Biochem. Biophys. Res. Commun. 280:1101-1106(2001).  
RN [2]  
RP SEQUENCE FROM N.A.  
RX MEDLINE=21092761; PubMed=1162673;  
RA Shukunami C., Oshima Y., Hiraki Y.;  
RT "Molecular cloning of tenomodulin, a novel chondromodulin-I related  
RT gene";  
RL Biochem. Biophys. Res. Commun. 280:1323-1327(2001).  
RN [3]  
RP SEQUENCE FROM N.A.  
RA TISSUE-Skeletal muscle;  
RA Cros N., Tratchenko A.V., Leclerc L., Leger J.J., Marini J.-F.,  
RA Dechesne C.A.;  
RT "Gene expression alterations revealed by suppression subtractive  
RT hybridization in rat soleus muscle disease atrophy";  
RL Submitted (OCT-1999) to the EMBL/GenBank/DBJ databases.  
RN [4]  
RP SEQUENCE OF 61-307 FROM N.A.  
RA Lawlor S.;  
RT Submitted (SEP-1999) to the EMBL/GenBank/DBJ databases.  
RL  
CC - FUNCTION: May be an angiogenesis inhibitor.  
CC - SUBCELLULAR LOCATION: Type II membrane protein (Potential).  
CC - SIMILARITY: Belongs to the chondromodulin-I family.  
CC - SIMILARITY: Contains 1 BRICHOS domain.  
-----  
CC This SWISS-PROT entry is copyright. It is produced through a collaboration  
CC between the Swiss Institute of Bioinformatics and the EMBL outstation -  
CC the European Bioinformatics Institute. There are no restrictions on its  
CC use by non-profit institutions as long as its content is in no way  
CC modified and this statement is not removed. Usage by and for commercial  
CC entities requires a license agreement (See <http://www.ebi.ac.uk/announcements>  
CC or send an email to [license@ebi.ac.uk](mailto:license@ebi.ac.uk)).  
-----  
CC EMBL; AB055421; BAB21756.1; -  
DR EMBL; AF234259; AAG9144.1; -  
DR EMBL; AF191770; AAG28395.1; -





```

QY 1 MAKNPENCEDCHILNAEAFKSKKICKSLKICGLVFGILATLILVFWGSKHFWPEVPRK 60
DB 1 MAKNPENCCEGCHILNAEALSKKICKSLKICGLVFGILATLILVFWGSKHFWPEVSKK 60
QY 61 AYDMEHTFYSNGEKKIKYMEIDPVTRETFRSNGTDETLVHDFKNGYTGIVGLOK 120
DB 61 TYDMEHTFYSNGEKKIKYMEIDPVTRETFRSNGTDETLVHDFKNGYTGIVGLOK 120
QY 121 FIKTQIKVIPFSPBEEIDENEITTTFFEQSVIWPAPKEPIENRDFLNSKILEICDN 180
DB 121 FIKTQIKVIPFSPBEEIDENEITTTFFEQSVIWPAPKEPIENRDFLNSKILEICDN 180
QY 181 VTMWNPPTLISVSELQDFEEGEDLHPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
DB 181 VTMWNPPTLISVSELQDFEEGEDLHPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
QY 241 BELPINDYENGIEFDPMLDERGYCCICRGRNRCRVCBPLLGYPYPCYGGGVIC 300
DB 241 EDLPNDYENGIEFDPMLDERGYCCICRGRNRCRVCBPLLGYPYPCYGGGVIC 300
QY 301 RVIMPCNMWVARMGRV 317
DB 301 RVIMPCNMWVARMGRV 317

RESULT 3
ID TNDM RAT STANDARD; PRT; 317 AA.
AC Q9ESC2;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 15-MAR-2004 (Rel. 43, Last annotation update)
DE Tenomodulin (Tm) (Tm) (Chondromodulin-I like protein) (ChMLL)
DE (ChMLL) (Myodulin).
GN TNDM OR CHML.
OS Rattus norvegicus (Rat).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OC NCBI_TaxId=10116;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=Sprague-Dawley; TISSUE=Skeletal muscle;
RX MEDLINE=21092726; PubMed=11162640;
RA Yamana K., Wada H., Takahashi Y., Sato H., Kasahara Y., Kiyoki M.;
RT "Molecular cloning and characterization of ChML, a novel membrane
RT molecule similar to chondromodulin-1."
RL Biochem. Biophys. Res. Commun. 280:1101-1106(2001).
RN [2]
RP SEQUENCE FROM N.A.
RA Crous N., Tkatchenko A.V., Leger J.J., Martin J.-F.,
RA Dechene C.A.;
RT "Gene expression alterations revealed by suppression subtractive
RT hybridization in rat soleus muscle disuse atrophy."
RL Submitted (Oct-1999) to the EMBL/GenBank/DBJ databases.
CC -1- FUNCTION: May be an angiogenesis inhibitor.
CC -1- SUBCELLULAR LOCATION: Type II membrane protein (Potential).
CC -1- SIMILARITY: Belongs to the chondromodulin-I family.
CC -1- SIMILARITY: Contains 1 BRICHOS domain.
CC
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC the European Bioinformatics Institute. There are no restrictions on its
CC use by non-profit institutions as long as its content is in no way
CC modified and this statement is not removed. Usage by and for commercial
CC entities requires a license agreement (See http://www.ebi.ac.uk/announcements/
CC or send an email to licenses@ebi.ac.uk).
CC
CC EMBL; AF191769; AAC28394.1; -
CC EMBL; AB055423; BAB21758.1; -
CC InterPro; IPR007084; BRICHOS.
CC Pfam; PF04089; BRICHOS.1.
CC PROSITE; PS50869; BRICHOS.1.
CC Glycoprotein; Transmembrane; Signal-anchor.

```

```

FT DOMAIN 1 30 CYTOPLASMIC (POTENTIAL).
FT TRANSMEM 31 50 SIGNAL-ANCHOR (TYPE-II MEMBRANE PROTEIN)
FT 51 317 (POTENTIAL).
FT DOMAIN 51 186 EXTRACELLULAR (POTENTIAL).
FT CARBOHYD 93 94 BRICHOS. (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 94 94 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 180 180 N-LINKED (GLCNAC. . .) (POTENTIAL).
SQ SEQUENCE 317 AA; 36971 MW; 8C0328825F915D88 CRC64;

Query Match 95.0%; Score 1657; DB 1; Length 317;
Best Local Similarity 94.6%; Pred. No. 9.1e-116;
Matches 300; Conservative 7; Mismatches 10; Indels 0; Gaps 0;

QY 1 MAKNPENCEDCHILNAEAFKSKKICKSLKICGLVFGILATLILVFWGSKHFWPEVPRK 60
DB 1 MAKNPENCCEGCHILNAEALSKKICKSLKICGLVFGILATLILVFWGSKHFWPEVSKK 60
QY 61 AYDMEHTFYSNGEKKIKYMEIDPVTRETFRSNGTDETLVHDFKNGYTGIVGLOK 120
DB 61 TYDMEHTFYSNGEKKIKYMEIDPVTRETFRSNGTDETLVHDFKNGYTGIVGLOK 120
QY 121 FIKTQIKVIPFSPBEEIDENEITTTFFEQSVIWPAPKEPIENRDFLNSKILEICDN 180
DB 121 FIKTQIKVIPFSPBEEIDENEITTTFFEQSVIWPAPKEPIENRDFLNSKILEICDN 180
QY 181 VTMWNPPTLISVSELQDFEEGEDLHPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
DB 181 VTMWNPPTLISVSELQDFEEGEDLHPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
QY 241 BELPINDYENGIEFDPMLDERGYCCICRGRNRCRVCBPLLGYPYPCYGGGVIC 300
DB 241 EDLPNDYENGIEFDPMLDERGYCCICRGRNRCRVCBPLLGYPYPCYGGGVIC 300
QY 301 RVIMPCNMWVARMGRV 317
DB 301 RVIMPCNMWVARMGRV 317

RESULT 4
ID CHML CHICK STANDARD; PRT; 347 AA.
AC Q9PUB8; Q9Y163;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 15-MAR-2004 (Rel. 43, Last annotation update)
DE Chondromodulin-I precursor (Chm-I) (Leukocyte cell-derived chemotaxin
DE 1) (contains: Chondrosulfatant protein (Ch-SP)).
GN LECT1 OR CHML.
OS Gallus gallus (Chicken).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Archosauria; Aves; Neognathae; Galliformes; Phasianidae; Phasianinae;
OC Gallus.
OC NCBI_TaxId=9031;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=White leghorn; TISSUE=Petal sternum;
RX MEDLINE=99379620; PubMed=1042551;
RA Shukunant C., Yamamoto S., Tarabe T., Hiraki Y.;
RT "Generation of multiple transcripts from the chicken chondromodulin-I
RT gene and their expression during embryonic development."
RL FBS Lett. 456:165-170(1999).
CC -1- FUNCTION: Bifunctional growth regulator. May contribute to the
CC rapid growth of cartilage and vascular invasion prior to the
CC replacement of cartilage by bone during endochondral bone
CC development (By similarity).

```



```

DB 189 ILSECDGLPIFWLKP--MYPKELPRERREVSSASTSTRPHSPRGNAFGLSLNRR 246
QY 234 HAQASBEEL-PINDYTE---NGIEFDPMLDERGYCCICRGRNRCRVCCELLGYPPY 289
DB 247 PSVQDDEEPNPNPNPHQOGESMTFDPRLDHGICICRGRSYTHCKICEPLGYYFW 306
QY 290 PNYCGGRCVIGVIMPCNMVYARMLGRV 317
DB 307 PNYCGGRCVIGVIMPCNMVYARMLGRV 334

RESULT 6
CHM1_MOUSE
ID CHM1_MOUSE STANDARD; PRT; 334 AA.
AC 0921F6; 09CXU5;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 15-MAR-2004 (Rel. 43, Last annotation update)
DE Chondromodulin-1 precursor (CHM-1) (Leukocyte cell-derived chemotaxin
1) [Contains: Chondrosurfactant protein (CH-SP)].
GN LECT1 OR CHM1.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxId=10090;
RN [1]
RP SEQUENCE FROM N.A.
RA Hiraki Y., Shukunani C., Inoue H., Suzuki F.;
RL Submitted (DEC-1995) to the EMBL/GenBank/DBJ databases.
RN [2]
RP SEQUENCE FROM N.A.
RA STRAIN=C57BL/6J; TISSUE=Embryonic head;
RX MEDLINE=21085660; PubMed=11217851;
RA Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA Atakawa T., Hara A., Fukunishi Y., Konno H., Adachi T., Fukuda S.,
RA Atzawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamataka I.,
RA Saito T., Okazaki Y., Gotohori T., Bono H., Kasukawa T., Saito R.,
RA Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA Kiehl P., Lewis S., Matsuo Y., Nikaido I., Pesole G., Quackenbush J.,
RA Schiraldi L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Beffelli D., Bojunga N., Carinci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli U., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyooka K., Wang K.H., Weitz C., Whitaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohseuk K.,
RA Hayashizaki Y.;
RA "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
CC -1- FUNCTION: Bifunctional growth regulator that stimulates the growth
of cultured chondrocytes in the presence of basic fibroblast
growth factor (FGF) but inhibits the growth of cultured vascular
endothelial cells. May contribute to the rapid growth of cartilage
and vascular invasion prior to the replacement of cartilage by
bone during endochondral bone development. (By similarity)
CC -1- SUBCELLULAR LOCATION: Cytoplasmic and secreted. Accumulated in the
inter-territorial matrix of cartilage. (By similarity)
CC -1- PM: After cleavage, the post-translational modified CHM-1 is
secreted as a glycoprotein (By similarity).
CC -1- SIMILARITY: Belongs to the chondromodulin-1 family.
CC -1- SIMILARITY: Contains 1 BRCHOS domain.
CC -----
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC the European Bioinformatics Institute. There are no restrictions on its
CC use by non-profit institutions as long as its content is in no way
CC modified and this statement is not removed. Usage by and for commercial
CC entities requires a license agreement (See http://www.isb-sib.ch/announce/)

```

```

CC or send an email to license@isb-sib.ch.
CC -----
CC DR EMBL; U43509; AAD00027.1; -.
CC DR EMBL; AK013975; BAB29095.1; -.
CC DR MGD; MGI:3341171; LECT1.
CC DR InterPro; IPR007084; BRCHOS.
CC DR Pfam; PF04089; BRCHOS; 1.
CC DR ProSite; PS50869; BRCHOS; 1.
CC DR Glycoprotein; Transmembrane; Cleavage on pair of basic residues;
CC Chondrogenesis.
CC CHAIN 1 210
CC PROPEP 211 214
CC CHAIN 215 334
CC TRANSMEM 46 66
CC DOMAIN 105 201
CC DISULFID 282 286
CC CARBOHYD 283 323
CC CONFLICT 243 243
CC CONFLICT 108 109
CC CONFLICT 163 163
CC CONFLICT 207 207
CC SEQUENCE 334 AA; 37252 MW; DAFDEC43CTD745D6 CRC64;
Query Match 29.7%; Score 518; DB 1; Length 334;
Best Local Similarity 36.3%; Pred. No. 6.4e-32;
Matches 119; Conservative 65; Mismatches 120; Indels 24; Gaps 11;
QY 6 PENCEDCHILNAEAKSKICKS-----LKICGLVFGIALLTLVFW--GSKHFWDEV 57
DB 15 PEDVEFC---SPFATVTVKPSGSPTRLKGVAV--LISGAVLLFGALGAFYFKGN 69
QY 58 PKKADMEHTFYSNKKKIMELDPVTRTEIPSGNQTDETLFHPDKNGYTGIFYGL 117
DB 70 DNHIVYVSSINIKQDGSNEIDAVNNLTFPKKSGAKALEVNDKNGITGRFAGG 129
QY 118 OKCFIKYIKV-IPFP-SEPEEIDENE--ITTFPQSYIWPAPKPIENRPLKNSK 173
DB 130 EECYIKAVKRIPEVGVTVKQISLELGKIMPNVYENSLIYVAVDQPVKDSFL-SSK 188
QY 174 ILEIDNVTWYINFTLISVSELDPEEFGEDLHPANEKGIENEGWVYPOVKEVTR 233
DB 189 ILELGDGLPIFWLKP--MYPKELQERREVANSAPSTRPHSPRGNAFGLSLNCTR 246
QY 234 -HARQASBEEL-PINDYTE---NGIEFDPMLDERGYCCICRGRNRCRVCCELLGYPPY 289
DB 247 PSVQDDEEPNPNPNPHQOGESMTFDPRLDHGICICRGRSYTHCKICEPLGYYFW 306
QY 290 PNYCGGRCVIGVIMPCNMVYARMLGRV 317
DB 307 PNYCGGRCVIGVIMPCNMVYARMLGRV 334

RESULT 7
CHM1_HUMAN
ID CHM1_HUMAN STANDARD; PRT; 334 AA.
AC 075829; 09UM18;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 15-MAR-2004 (Rel. 43, Last annotation update)
DE Chondromodulin-1 precursor (CHM-1) (Leukocyte cell-derived chemotaxin
1) [Contains: Chondrosurfactant protein (CH-SP)].
GN LECT1 OR CHM1.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Homiidae; Homo.
OX NCBI_TaxId=9606;
RN [1]
RP SEQUENCE FROM N.A.
RA TISSUE=Chondrosarcoma;
RX MEDLINE=98401184; PubMed=9731231;
RA Shukunani C., Hiraki Y.;
RA "Expression of cartilage-specific functional matrix chondromodulin-1

```

RT mRNA in rabbit growth plate chondrocytes and its responsiveness to  
RT growth stimuli in vitro.";  
RT Biochem. Biophys. Res. Commun. 249:885-890(1998).  
RN [2]  
RN SEQUENCE FROM N.A.  
RC TISSUE=Chondrosarcoma;  
RX MEDLINE=99203485; PubMed=10103019;  
RA Hiraki Y., Mitsui K., Endo N., Takahashi K., Hayami T., Inoue H.,  
RA Shukunami C., Tokunaga K., Kono T., Yamada M., Takahashi H.E.,  
RA Kondo J.;  
RT "Molecular cloning of human chondromodulin-I, a cartilage-derived  
RT growth modulating factor, and its expression in Chinese hamster ovary  
RT cells.";  
RT Eur. J. Biochem. 260:863-878(1999).  
RN [3]  
RN SEQUENCE OF 1-20 FROM N.A.  
RN Ozono K.;  
RN Submitted (DEC-1998) to the EMBL/GenBank/DBJ databases.  
RN [4]  
RN SEQUENCE OF 215-334 FROM N.A.  
RX MEDLINE=20035732; PubMed=10570955;  
RA Hayami T., Shukunami C., Mitsui K., Endo N., Tokunaga K., Kondo J.,  
RA Takahashi H.E., Hiraki Y.;  
RT "Specific loss of chondromodulin-I gene expression in chondrosarcoma  
RT and the suppression of tumor angiogenesis and growth by its  
RT recombinant protein in vivo.";  
RT FEBS Lett. 458:436-440(1999).  
RN [5]  
RN REVIEW.  
RX MEDLINE=20367654; PubMed=10912526;  
RA Hiraki Y., Shukunami C.;  
RT "Chondromodulin-I as a novel cartilage-specific growth-modulating  
RT factor.";  
RT Pediatr. Nephrol. 14:602-605(2000).  
CC -1- FUNCTION: Bifunctional growth regulator that stimulates the growth  
CC of cultured chondrocytes in the presence of basic fibroblast  
CC growth factor (FGF) but inhibits the growth of cultured vascular  
CC endothelial cells. May contribute to the rapid growth of cartilage  
CC and vascular invasion prior to the replacement of cartilage by  
CC bone during endochondral bone development.  
CC -1- SUBCELLULAR LOCATION: Cytoplasmic and secreted. Accumulated in the  
CC inter-territorial matrix of cartilage.  
CC -1- TISSUE SPECIFICITY: Cartilage specific. Weakly expressed in  
CC chondrosarcoma.  
CC -1- DEVELOPMENTAL STAGE: Expressed at 9 weeks in developing  
CC cartilaginous bone rudiments.  
CC -1- PTM: After cleavage, the post-translationally modified Chm-I is  
CC secreted as a glycoprotein.  
CC -1- SIMILARITY: Belongs to the chondromodulin-I family.  
CC -1- SIMILARITY: Contains 1 BRICHOS domain.  
CC -----  
CC This SWISS-PROT entry is copyright. It is produced through a collaboration  
CC between the Swiss Institute of Bioinformatics and the EMBL outstation -  
CC the European Bioinformatics Institute. There are no restrictions on its  
CC use by non-profit institutions as long as its content is in no way  
CC modified and this statement is not removed. Usage by and for commercial  
CC entities requires a license agreement (see <http://www.isb-sib.ch/announce/>  
CC or send an email to [license@sib-sib.ch](mailto:license@sib-sib.ch)).  
CC -----  
CC EMBL; AB006000; BAA33443.1; -  
CC EMBL; AF050147; AAC98971.1; -  
CC EMBL; AB021290; BAA77384.1; -  
CC EMBL; AB005999; BAA66262.1; -  
CC Genew; HGNC:17005; LECT1.  
CC MIM; 605147; -  
DR GO; GO:0008151; P:cell growth and/or maintenance; TAS.  
DR GO; GO:0006029; P:proteoglycan metabolism; TAS.  
DR GO; GO:0001501; P:skeletal development; TAS.  
DR InterPro; IPR007084; BRICHOS.  
DR Pfam; PF04089; BRICHOS; 1.  
DR PROSITE; PS50869; BRICHOS; 1.  
KM Glycoprotein; Transmembrane; Cleavage on pair of basic residues;  
KM Chondrogenesis.

FT CHAIN 1 210 CHONDROREFRACTANT PROTEIN  
FT PROPEP 211 214 (BY SIMILARITY).  
FT CHAIN 215 334 POTENTIAL.  
FT TRANSMEM 45 65 CHONDROMODULIN-I.  
FT DOMAIN 104 201 POTENTIAL.  
FT DISULFID 282 286 BRICHOS.  
FT DISULFID 283 323 BY SIMILARITY.  
FT CARBOHYD 243 243 BY SIMILARITY.  
SQ SEQUENCE 334 AA; 37102 MW; 9E239311F9D4F85 CRC64;  
Query Match 29.6%; Score 516; DB 1; Length 334;  
Best Local Similarity 35.2%; Pred. No. 9.1e-32;  
Matches 118; Conservative 63; Mismatches 116; Indels 38; Gaps 12;  
QY 6 PENCECHILNAPF-----KSKICKSLKIGLFGILALILYLFW--GSKHWPPEVP 58  
DB 15 PDVVEFC---SPRAYTLTVKSPSPALLKVAAY--LLISAVLLFGAIGAFYFWKGS 69  
QY 59 KKAYDHEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETVHDFKNGYTGIFVGLQ 118  
DB 70 SHLYNHHTMSNGKIQDSMETIDAGNNLETRMSGABEALAVNDFOGIIIGIRPAGE 129  
QY 119 KCFIKYQIVY-IPER-SEPEEEDID--ENEITTFEESQVIVWPAKTEINRDFIKSK 173  
DB 130 KCYIKAQVAPRIPEVAVTKOSISSKLECKIMPVXEENSLIYVAVDQPKONSFL-SSK 188  
QY 174 ILEICGNVMTWVNIPTLISVSE-ODEFEEDGLHPANEKK-----GIEQNEQM 222  
DB 189 VLEICGDLPIFWIKPYY--PKELQREKRVYAKIYPTTKRHSGSRNPAGRLNNE-- 244  
QY 223 VPDVVEKTRHARQASBEELPINDYENGIEFDPMLDERGYCCICGRGNYRCRVCEP 282  
DB 245 TRSVQEDS-----QAFNPDPNPHQEGESMTFDEPLDHEGICIECRSSYTHCKICEP 299  
QY 283 LGGYEPPYCYCGGRVICVIMPCNMVYARMGRV 317  
DB 300 LGGYEPPYNYOGCRACRACVIMPCSWVARILGMV 334  
RESULT 8  
CHM1\_BOVIN STANDARD; PRT; 335 AA.  
ID P17404; P23590;  
AC 01-AUG-1990 (Rel. 15, Created)  
DT 01-NOV-1991 (Rel. 20, Last sequence update)  
DT 15-MAR-2004 (Rel. 43, Last annotation update)  
DE Chondromodulin-I precursor (Chm-I) (Leukocyte cell-derived chemotaxin  
DE 1) (Small cartilage-derived glycoprotein) (SCGP) [contains:  
DE Chondrosurfactant protein (CH-SP)].  
GN LECT1 OR CHM1.  
OS Bos taurus (Bovine).  
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
OC Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae;  
OC Bovidae; Bovinae; Bos.  
OX NCBI\_TaxID=9913;  
RN [1]  
RN SEQUENCE FROM N.A. AND SEQUENCE OF 215-241 AND 297-317.  
RC TISSUE=fetal epiphyseal cartilage;  
RX MEDLINE=9122210; PubMed=1709014;  
RA Hiraki Y., Tanaka H., Inoue H., Kondo J., Kamizono A., Suzuki F.;  
RT "Molecular cloning of a new class of cartilage-specific matrix,  
RT chondromodulin-I, which stimulates growth of cultured chondrocytes.";  
RT Biochem. Biophys. Res. Commun. 175:971-977(1991).  
RN [2]  
RN SEQUENCE OF 215-335, VARIANT MET-221, AND GLYCOSYLATION.  
RC TISSUE=Nasal cartilage;  
RX MEDLINE=90277643; PubMed=2351661;  
RA Namee P.J., Treep J.T., Young C.N.;  
RT "An 18-kDa glycoprotein from bovine nasal cartilage. Isolation and  
RT primary structure of small, cartilage-derived glycoprotein.";  
RL J. Biol. Chem. 265:9628-9633(1990).  
RN [3]

RP ERRATUM.  
RA Name P.J., Treep J.T., Young C.N.;  
RL J. Biol. Chem. 265:22056-22056(1990).  
RN [4]  
RP PROCESSING.  
RX MEDLINE=21316541; PubMed=11323410;  
RA Azizan A., Holaday N., Name P.O.;  
RT Post-translational processing of bovine chondromodulin-I.;  
RL J. Biol. Chem. 276:23632-23638 (2001).  
RN [5]  
RP REVIEW.  
RX MEDLINE=20367654; PubMed=10912526;  
RA Hiraki Y., Shukunami C.;  
RT "Chondromodulin-I as a novel cartilage-specific growth-modulating factor."  
RL Pediatr. Nephrol. 14:602-605(2000).  
CC -1- FUNCTION: Bifunctional growth regulator that stimulates the growth of cultured chondrocytes in the presence of basic fibroblast growth factor (FGF) but inhibits the growth of cultured vascular endothelial cells. May contribute to the rapid growth of cartilage and vascular invasion prior to the replacement of cartilage by bone during endochondral bone development.  
CC -1- SUBCELLULAR LOCATION: Cytoplasmic and secreted. Accumulated in the inter-territorial matrix of cartilage.  
CC -1- TISSUE SPECIFICITY: Nasal and articular cartilage, and fetal epiphysis.  
CC -1- PTM: After cleavage, the post-translationally modified Chm-I is secreted as a glycoprotein.  
CC -1- PTM: Two other smaller nonglycosylated chondromodulin forms (9 kDa and 7kDa) are found either in developing articular cartilage or in chondrocytes. The 9 kDa form could be processed by an extracellular matrix-associated protease as a metalloproteinase and the 7 kDa form could be processed intracellularly.  
CC -1- SIMILARITY: Belongs to the chondromodulin-I family.  
CC -1- SIMILARITY: Contains 1 BRICHOS domain.  
CC This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation - the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See <http://www.isb-sib.ch/announce/> or send an email to [license@isb-sib.ch](mailto:license@isb-sib.ch)).  
CC -----  
DR EMBL; M65081; AAA30445.1; -.  
DR PIR; J0569; J0569.  
DR InterPro; IPR007084; BRICHOS.  
DR Pfam; PF04089; BRICHOS.1.  
DR PROSITE; PS00869; BRICHOS.1.  
KW Glycoprotein; Transmembrane; Cleavage on pair of basic residues; Chondrogenesis.  
FT CHAIN 1 210 CHONDROSURFACTANT PROTEIN (POTENTIAL).  
FT PROPEP 211 214 POTENTIAL.  
FT CHAIN 215 335 CHONDROMODULIN-I.  
FT TRANSMEM 45 65 POTENTIAL.  
FT DOMAIN 104 201 BRICHOS.  
FT DISULFID 283 287  
FT DISULFID 284 324  
FT CARBOHYD 223 223  
FT CARBOHYD 236 236  
FT CARBOHYD 244 244  
FT CARBOHYD 221 221  
FT VARIANT 221 221  
FT VARIANT 222 222  
FT VARIANT 223 223  
FT VARIANT 224 224  
FT VARIANT 225 225  
FT VARIANT 226 226  
FT VARIANT 227 227  
FT VARIANT 228 228  
FT VARIANT 229 229  
FT VARIANT 230 230  
FT VARIANT 231 231  
FT VARIANT 232 232  
FT VARIANT 233 233  
FT VARIANT 234 234  
FT VARIANT 235 235  
FT VARIANT 236 236  
FT VARIANT 237 237  
FT VARIANT 238 238  
FT VARIANT 239 239  
FT VARIANT 240 240  
FT VARIANT 241 241  
FT VARIANT 242 242  
FT VARIANT 243 243  
FT VARIANT 244 244  
FT VARIANT 245 245  
FT VARIANT 246 246  
FT VARIANT 247 247  
FT VARIANT 248 248  
FT VARIANT 249 249  
FT VARIANT 250 250  
FT VARIANT 251 251  
FT VARIANT 252 252  
FT VARIANT 253 253  
FT VARIANT 254 254  
FT VARIANT 255 255  
FT VARIANT 256 256  
FT VARIANT 257 257  
FT VARIANT 258 258  
FT VARIANT 259 259  
FT VARIANT 260 260  
FT VARIANT 261 261  
FT VARIANT 262 262  
FT VARIANT 263 263  
FT VARIANT 264 264  
FT VARIANT 265 265  
FT VARIANT 266 266  
FT VARIANT 267 267  
FT VARIANT 268 268  
FT VARIANT 269 269  
FT VARIANT 270 270  
FT VARIANT 271 271  
FT VARIANT 272 272  
FT VARIANT 273 273  
FT VARIANT 274 274  
FT VARIANT 275 275  
FT VARIANT 276 276  
FT VARIANT 277 277  
FT VARIANT 278 278  
FT VARIANT 279 279  
FT VARIANT 280 280  
FT VARIANT 281 281  
FT VARIANT 282 282  
FT VARIANT 283 283  
FT VARIANT 284 284  
FT VARIANT 285 285  
FT VARIANT 286 286  
FT VARIANT 287 287  
FT VARIANT 288 288  
FT VARIANT 289 289  
FT VARIANT 290 290  
FT VARIANT 291 291  
FT VARIANT 292 292  
FT VARIANT 293 293  
FT VARIANT 294 294  
FT VARIANT 295 295  
FT VARIANT 296 296  
FT VARIANT 297 297  
FT VARIANT 298 298  
FT VARIANT 299 299  
FT VARIANT 300 300  
FT VARIANT 301 301  
FT VARIANT 302 302  
FT VARIANT 303 303  
FT VARIANT 304 304  
FT VARIANT 305 305  
FT VARIANT 306 306  
FT VARIANT 307 307  
FT VARIANT 308 308  
FT VARIANT 309 309  
FT VARIANT 310 310  
FT VARIANT 311 311  
FT VARIANT 312 312  
FT VARIANT 313 313  
FT VARIANT 314 314  
FT VARIANT 315 315  
FT VARIANT 316 316  
FT VARIANT 317 317  
FT VARIANT 318 318  
FT VARIANT 319 319  
FT VARIANT 320 320  
FT VARIANT 321 321  
FT VARIANT 322 322  
FT VARIANT 323 323  
FT VARIANT 324 324  
FT VARIANT 325 325  
FT VARIANT 326 326  
FT VARIANT 327 327  
FT VARIANT 328 328  
FT VARIANT 329 329  
FT VARIANT 330 330  
FT VARIANT 331 331  
FT VARIANT 332 332  
FT VARIANT 333 333  
FT VARIANT 334 334  
FT VARIANT 335 335  
FT VARIANT 336 336  
FT VARIANT 337 337  
FT VARIANT 338 338  
FT VARIANT 339 339  
FT VARIANT 340 340  
FT VARIANT 341 341  
FT VARIANT 342 342  
FT VARIANT 343 343  
FT VARIANT 344 344  
FT VARIANT 345 345  
FT VARIANT 346 346  
FT VARIANT 347 347  
FT VARIANT 348 348  
FT VARIANT 349 349  
FT VARIANT 350 350  
FT VARIANT 351 351  
FT VARIANT 352 352  
FT VARIANT 353 353  
FT VARIANT 354 354  
FT VARIANT 355 355  
FT VARIANT 356 356  
FT VARIANT 357 357  
FT VARIANT 358 358  
FT VARIANT 359 359  
FT VARIANT 360 360  
FT VARIANT 361 361  
FT VARIANT 362 362  
FT VARIANT 363 363  
FT VARIANT 364 364  
FT VARIANT 365 365  
FT VARIANT 366 366  
FT VARIANT 367 367  
FT VARIANT 368 368  
FT VARIANT 369 369  
FT VARIANT 370 370  
FT VARIANT 371 371  
FT VARIANT 372 372  
FT VARIANT 373 373  
FT VARIANT 374 374  
FT VARIANT 375 375  
FT VARIANT 376 376  
FT VARIANT 377 377  
FT VARIANT 378 378  
FT VARIANT 379 379  
FT VARIANT 380 380  
FT VARIANT 381 381  
FT VARIANT 382 382  
FT VARIANT 383 383  
FT VARIANT 384 384  
FT VARIANT 385 385  
FT VARIANT 386 386  
FT VARIANT 387 387  
FT VARIANT 388 388  
FT VARIANT 389 389  
FT VARIANT 390 390  
FT VARIANT 391 391  
FT VARIANT 392 392  
FT VARIANT 393 393  
FT VARIANT 394 394  
FT VARIANT 395 395  
FT VARIANT 396 396  
FT VARIANT 397 397  
FT VARIANT 398 398  
FT VARIANT 399 399  
FT VARIANT 400 400  
FT VARIANT 401 401  
FT VARIANT 402 402  
FT VARIANT 403 403  
FT VARIANT 404 404  
FT VARIANT 405 405  
FT VARIANT 406 406  
FT VARIANT 407 407  
FT VARIANT 408 408  
FT VARIANT 409 409  
FT VARIANT 410 410  
FT VARIANT 411 411  
FT VARIANT 412 412  
FT VARIANT 413 413  
FT VARIANT 414 414  
FT VARIANT 415 415  
FT VARIANT 416 416  
FT VARIANT 417 417  
FT VARIANT 418 418  
FT VARIANT 419 419  
FT VARIANT 420 420  
FT VARIANT 421 421  
FT VARIANT 422 422  
FT VARIANT 423 423  
FT VARIANT 424 424  
FT VARIANT 425 425  
FT VARIANT 426 426  
FT VARIANT 427 427  
FT VARIANT 428 428  
FT VARIANT 429 429  
FT VARIANT 430 430  
FT VARIANT 431 431  
FT VARIANT 432 432  
FT VARIANT 433 433  
FT VARIANT 434 434  
FT VARIANT 435 435  
FT VARIANT 436 436  
FT VARIANT 437 437  
FT VARIANT 438 438  
FT VARIANT 439 439  
FT VARIANT 440 440  
FT VARIANT 441 441  
FT VARIANT 442 442  
FT VARIANT 443 443  
FT VARIANT 444 444  
FT VARIANT 445 445  
FT VARIANT 446 446  
FT VARIANT 447 447  
FT VARIANT 448 448  
FT VARIANT 449 449  
FT VARIANT 450 450  
FT VARIANT 451 451  
FT VARIANT 452 452  
FT VARIANT 453 453  
FT VARIANT 454 454  
FT VARIANT 455 455  
FT VARIANT 456 456  
FT VARIANT 457 457  
FT VARIANT 458 458  
FT VARIANT 459 459  
FT VARIANT 460 460  
FT VARIANT 461 461  
FT VARIANT 462 462  
FT VARIANT 463 463  
FT VARIANT 464 464  
FT VARIANT 465 465  
FT VARIANT 466 466  
FT VARIANT 467 467  
FT VARIANT 468 468  
FT VARIANT 469 469  
FT VARIANT 470 470  
FT VARIANT 471 471  
FT VARIANT 472 472  
FT VARIANT 473 473  
FT VARIANT 474 474  
FT VARIANT 475 475  
FT VARIANT 476 476  
FT VARIANT 477 477  
FT VARIANT 478 478  
FT VARIANT 479 479  
FT VARIANT 480 480  
FT VARIANT 481 481  
FT VARIANT 482 482  
FT VARIANT 483 483  
FT VARIANT 484 484  
FT VARIANT 485 485  
FT VARIANT 486 486  
FT VARIANT 487 487  
FT VARIANT 488 488  
FT VARIANT 489 489  
FT VARIANT 490 490  
FT VARIANT 491 491  
FT VARIANT 492 492  
FT VARIANT 493 493  
FT VARIANT 494 494  
FT VARIANT 495 495  
FT VARIANT 496 496  
FT VARIANT 497 497  
FT VARIANT 498 498  
FT VARIANT 499 499  
FT VARIANT 500 500  
FT VARIANT 501 501  
FT VARIANT 502 502  
FT VARIANT 503 503  
FT VARIANT 504 504  
FT VARIANT 505 505  
FT VARIANT 506 506  
FT VARIANT 507 507  
FT VARIANT 508 508  
FT VARIANT 509 509  
FT VARIANT 510 510  
FT VARIANT 511 511  
FT VARIANT 512 512  
FT VARIANT 513 513  
FT VARIANT 514 514  
FT VARIANT 515 515  
FT VARIANT 516 516  
FT VARIANT 517 517  
FT VARIANT 518 518  
FT VARIANT 519 519  
FT VARIANT 520 520  
FT VARIANT 521 521  
FT VARIANT 522 522  
FT VARIANT 523 523  
FT VARIANT 524 524  
FT VARIANT 525 525  
FT VARIANT 526 526  
FT VARIANT 527 527  
FT VARIANT 528 528  
FT VARIANT 529 529  
FT VARIANT 530 530  
FT VARIANT 531 531  
FT VARIANT 532 532  
FT VARIANT 533 533  
FT VARIANT 534 534  
FT VARIANT 535 535  
FT VARIANT 536 536  
FT VARIANT 537 537  
FT VARIANT 538 538  
FT VARIANT 539 539  
FT VARIANT 540 540  
FT VARIANT 541 541  
FT VARIANT 542 542  
FT VARIANT 543 543  
FT VARIANT 544 544  
FT VARIANT 545 545  
FT VARIANT 546 546  
FT VARIANT 547 547  
FT VARIANT 548 548  
FT VARIANT 549 549  
FT VARIANT 550 550  
FT VARIANT 551 551  
FT VARIANT 552 552  
FT VARIANT 553 553  
FT VARIANT 554 554  
FT VARIANT 555 555  
FT VARIANT 556 556  
FT VARIANT 557 557  
FT VARIANT 558 558  
FT VARIANT 559 559  
FT VARIANT 560 560  
FT VARIANT 561 561  
FT VARIANT 562 562  
FT VARIANT 563 563  
FT VARIANT 564 564  
FT VARIANT 565 565  
FT VARIANT 566 566  
FT VARIANT 567 567  
FT VARIANT 568 568  
FT VARIANT 569 569  
FT VARIANT 570 570  
FT VARIANT 571 571  
FT VARIANT 572 572  
FT VARIANT 573 573  
FT VARIANT 574 574  
FT VARIANT 575 575  
FT VARIANT 576 576  
FT VARIANT 577 577  
FT VARIANT 578 578  
FT VARIANT 579 579  
FT VARIANT 580 580  
FT VARIANT 581 581  
FT VARIANT 582 582  
FT VARIANT 583 583  
FT VARIANT 584 584  
FT VARIANT 585 585  
FT VARIANT 586 586  
FT VARIANT 587 587  
FT VARIANT 588 588  
FT VARIANT 589 589  
FT VARIANT 590 590  
FT VARIANT 591 591  
FT VARIANT 592 592  
FT VARIANT 593 593  
FT VARIANT 594 594  
FT VARIANT 595 595  
FT VARIANT 596 596  
FT VARIANT 597 597  
FT VARIANT 598 598  
FT VARIANT 599 599  
FT VARIANT 600 600  
FT VARIANT 601 601  
FT VARIANT 602 602  
FT VARIANT 603 603  
FT VARIANT 604 604  
FT VARIANT 605 605  
FT VARIANT 606 606  
FT VARIANT 607 607  
FT VARIANT 608 608  
FT VARIANT 609 609  
FT VARIANT 610 610  
FT VARIANT 611 611  
FT VARIANT 612 612  
FT VARIANT 613 613  
FT VARIANT 614 614  
FT VARIANT 615 615  
FT VARIANT 616 616  
FT VARIANT 617 617  
FT VARIANT 618 618  
FT VARIANT 619 619  
FT VARIANT 620 620  
FT VARIANT 621 621  
FT VARIANT 622 622  
FT VARIANT 623 623  
FT VARIANT 624 624  
FT VARIANT 625 625  
FT VARIANT 626 626  
FT VARIANT 627 627  
FT VARIANT 628 628  
FT VARIANT 629 629  
FT VARIANT 630 630  
FT VARIANT 631 631  
FT VARIANT 632 632  
FT VARIANT 633 633  
FT VARIANT 634 634  
FT VARIANT 635 635  
FT VARIANT 636 636  
FT VARIANT 637 637  
FT VARIANT 638 638  
FT VARIANT 639 639  
FT VARIANT 640 640  
FT VARIANT 641 641  
FT VARIANT 642 642  
FT VARIANT 643 643  
FT VARIANT 644 644  
FT VARIANT 645 645  
FT VARIANT 646 646  
FT VARIANT 647 647  
FT VARIANT 648 648  
FT VARIANT 649 649  
FT VARIANT 650 650  
FT VARIANT 651 651  
FT VARIANT 652 652  
FT VARIANT 653 653  
FT VARIANT 654 654  
FT VARIANT 655 655  
FT VARIANT 656 656  
FT VARIANT 657 657  
FT VARIANT 658 658  
FT VARIANT 659 659  
FT VARIANT 660 660  
FT VARIANT 661 661  
FT VARIANT 662 662  
FT VARIANT 663 663  
FT VARIANT 664 664  
FT VARIANT 665 665  
FT VARIANT 666 666  
FT VARIANT 667 667  
FT VARIANT 668 668  
FT VARIANT 669 669  
FT VARIANT 670 670  
FT VARIANT 671 671  
FT VARIANT 672 672  
FT VARIANT 673 673  
FT VARIANT 674 674  
FT VARIANT 675 675  
FT VARIANT 676 676  
FT VARIANT 677 677  
FT VARIANT 678 678  
FT VARIANT 679 679  
FT VARIANT 680 680  
FT VARIANT 681 681  
FT VARIANT 682 682  
FT VARIANT 683 683  
FT VARIANT 684 684  
FT VARIANT 685 685  
FT VARIANT 686 686  
FT VARIANT 687 687  
FT VARIANT 688 688  
FT VARIANT 689 689  
FT VARIANT 690 690  
FT VARIANT 691 691  
FT VARIANT 692 692  
FT VARIANT 693 693  
FT VARIANT 694 694  
FT VARIANT 695 695  
FT VARIANT 696 696  
FT VARIANT 697 697  
FT VARIANT 698 698  
FT VARIANT 699 699  
FT VARIANT 700 700  
FT VARIANT 701 701  
FT VARIANT 702 702  
FT VARIANT 703 703  
FT VARIANT 704 704  
FT VARIANT 705 705  
FT VARIANT 706 706  
FT VARIANT 707 707  
FT VARIANT 708 708  
FT VARIANT 709 709  
FT VARIANT 710 710  
FT VARIANT 711 711  
FT VARIANT 712 712  
FT VARIANT 713 713  
FT VARIANT 714 714  
FT VARIANT 715 715  
FT VARIANT 716 716  
FT VARIANT 717 717  
FT VARIANT 718 718  
FT VARIANT 719 719  
FT VARIANT 720 720  
FT VARIANT 721 721  
FT VARIANT 722 722  
FT VARIANT 723 723  
FT VARIANT 724 724  
FT VARIANT 725 725  
FT VARIANT 726 726  
FT VARIANT 727 727  
FT VARIANT 728 728  
FT VARIANT 729 729  
FT VARIANT 730 730  
FT VARIANT 731 731  
FT VARIANT 732 732  
FT VARIANT 733 733  
FT VARIANT 734 734  
FT VARIANT 735 735  
FT VARIANT 736 736  
FT VARIANT 737 737  
FT VARIANT 738 738  
FT VARIANT 739 739  
FT VARIANT 740 740  
FT VARIANT 741 741  
FT VARIANT 742 742  
FT VARIANT 743 743  
FT VARIANT 744 744  
FT VARIANT 745 745  
FT VARIANT 746 746  
FT VARIANT 747 747  
FT VARIANT 748 748  
FT VARIANT 749 749  
FT VARIANT 750 750  
FT VARIANT 751 751  
FT VARIANT 752 752  
FT VARIANT 753 753  
FT VARIANT 754 754  
FT VARIANT 755 755  
FT VARIANT 756 756  
FT VARIANT 757 757  
FT VARIANT 758 758  
FT VARIANT 759 759  
FT VARIANT 760 760  
FT VARIANT 761 761  
FT VARIANT 762 762  
FT VARIANT 763 763  
FT VARIANT 764 764  
FT VARIANT 765 765  
FT VARIANT 766 766  
FT VARIANT 767 767  
FT VARIANT 768 768  
FT VARIANT 769 769  
FT VARIANT 770 770  
FT VARIANT 771 771  
FT VARIANT 772 772  
FT VARIANT 773 773  
FT VARIANT 774 774  
FT VARIANT 775 775  
FT VARIANT 776 776  
FT VARIANT 777 777  
FT VARIANT 778 778  
FT VARIANT 779 779  
FT VARIANT 780 780  
FT VARIANT 781 781  
FT VARIANT 782 782  
FT VARIANT 783 783  
FT VARIANT 784 784  
FT VARIANT 785 785  
FT VARIANT 786 786  
FT VARIANT 787 787  
FT VARIANT 788 788  
FT VARIANT 789 789  
FT VARIANT 790 790  
FT VARIANT 791 791  
FT VARIANT 792 792  
FT VARIANT 793 793  
FT VARIANT 794 794  
FT VARIANT 795 795  
FT VARIANT 796 796  
FT VARIANT 797 797  
FT VARIANT 798 798  
FT VARIANT 799 799  
FT VARIANT 800 800  
FT VARIANT 801 801  
FT VARIANT 802 802  
FT VARIANT 803 803  
FT VARIANT 804 804  
FT VARIANT 805 805  
FT VARIANT 806 806  
FT VARIANT 807 807  
FT VARIANT 808 808  
FT VARIANT 809 809  
FT VARIANT 810 810  
FT VARIANT 811 811  
FT VARIANT 812 812  
FT VARIANT 813 813  
FT VARIANT 814 814  
FT VARIANT 815 815  
FT VARIANT 816 816  
FT VARIANT 817 817  
FT VARIANT 818 818  
FT VARIANT 819 819  
FT VARIANT 820 820  
FT VARIANT 821 821  
FT VARIANT 822 822  
FT VARIANT 823 823  
FT VARIANT 824 824  
FT VARIANT 825 825  
FT VARIANT 826 826  
FT VARIANT 827 827  
FT VARIANT 828 828  
FT VARIANT 829 829  
FT VARIANT 830 830  
FT VARIANT 831 831  
FT VARIANT 832 832  
FT VARIANT 833 833  
FT VARIANT 834 834  
FT VARIANT 835 835  
FT VARIANT 836 836  
FT VARIANT 837 837  
FT VARIANT 838 838  
FT VARIANT 839 839  
FT VARIANT 840 840  
FT VARIANT 841 841  
FT VARIANT 842 842  
FT VARIANT 843 843  
FT VARIANT 844 844  
FT VARIANT 845 845  
FT VARIANT 846 846  
FT VARIANT 847 847  
FT VARIANT 848 848  
FT VARIANT 849 849  
FT VARIANT 850 850  
FT VARIANT 851 851  
FT VARIANT 852 852  
FT VARIANT 853 853  
FT VARIANT 854 854  
FT VARIANT 855 855  
FT VARIANT 856 856  
FT VARIANT 857 857  
FT VARIANT 858 858  
FT VARIANT 859 859  
FT VARIANT 860 860  
FT VARIANT 861 861  
FT VARIANT 862 862  
FT VARIANT 863 863  
FT VARIANT 864 864  
FT VARIANT 865 865  
FT VARIANT 866 866  
FT VARIANT 867 867  
FT VARIANT 868 868  
FT VARIANT 869 869  
FT VARIANT 870 870  
FT VARIANT 871 871  
FT VARIANT 872 872  
FT VARIANT 873 873  
FT VARIANT 874 874  
FT VARIANT 875 875  
FT VARIANT 876 876  
FT VARIANT 877 877  
FT VARIANT 878 878  
FT VARIANT 879 879  
FT VARIANT 880 880  
FT VARIANT 881 881  
FT VARIANT 882 882  
FT VARIANT 883 883  
FT VARIANT 884 884  
FT VARIANT 885 885  
FT VARIANT 886 886  
FT VARIANT 887 887  
FT VARIANT 888 888  
FT VARIANT 889 889  
FT VARIANT 890 890  
FT VARIANT 891 891  
FT VARIANT 892 892  
FT VARIANT 893 893  
FT VARIANT 894 894  
FT VARIANT 895 895  
FT VARIANT 896 896  
FT VARIANT 897 897  
FT VARIANT 898 898  
FT VARIANT 899 899  
FT VARIANT 900 900  
FT VARIANT 901 901  
FT VARIANT 902 902  
FT VARIANT 903 903  
FT VARIANT 904 904  
FT VARIANT 905 905  
FT VARIANT 906 906  
FT VARIANT 907 907  
FT VARIANT 908 908  
FT VARIANT 909 909  
FT VARIANT 910 910  
FT VARIANT 911 911  
FT VARIANT 912 912  
FT VARIANT 913 913  
FT VARIANT 914 914  
FT VARIANT 915 915  
FT VARIANT 916 916  
FT VARIANT 917 917  
FT VARIANT 918 918  
FT VARIANT 919 919  
FT VARIANT 920 920  
FT VARIANT 921 921  
FT VARIANT 922 922  
FT VARIANT 923 923  
FT VARIANT 924 924  
FT VARIANT 925 925  
FT VARIANT 926 926  
FT VARIANT 927 927  
FT VARIANT 928 928  
FT VARIANT 929 929  
FT VARIANT 930 930  
FT VARIANT 931 931  
FT VARIANT 932 932  
FT VARIANT 933 933  
FT VARIANT 934 934  
FT VARIANT 935 935  
FT VARIANT 936 936  
FT VARIANT 937 937  
FT VARIANT 938 938  
FT VARIANT 939 939  
FT VARIANT 940 940  
FT VARIANT 941 941  
FT VARIANT 942 942  
FT VARIANT 943 943  
FT VARIANT 944 944  
FT VARIANT 945 945  
FT VARIANT 946 946  
FT VARIANT 947 947  
FT VARIANT 948 948  
FT VARIANT 949 949  
FT VARIANT 950 950  
FT VARIANT 951 951  
FT VARIANT 952 952  
FT VARIANT 953 953  
FT VARIANT 954 954  
FT VARIANT 955 955  
FT VARIANT 956 956  
FT VARIANT 957 957  
FT VARIANT 958 958  
FT VARIANT 959 959  
FT VARIANT 960 960  
FT VARIANT 961 961  
FT VARIANT 962 962  
FT VARIANT 963 963  
FT VARIANT 964 964  
FT VARIANT 965 965  
FT VARIANT 966 966  
FT VARIANT 967 967  
FT VARIANT 968 968  
FT VARIANT 969 969  
FT VARIANT 970 970  
FT VARIANT 971 971  
FT VARIANT 972 972  
FT VARIANT 973 973  
FT VARIANT 974 974  
FT VARIANT 975 975  
FT VARIANT 976 976  
FT VARIANT 977 977  
FT VARIANT 978 978  
FT VARIANT 979 979  
FT VARIANT 980 980  
FT VARIANT 981 981  
FT VARIANT 982 982  
FT VARIANT 983 983  
FT VARIANT 984 984  
FT VARIANT 985 985  
FT VARIANT 986 986  
FT VARIANT 987 987  
FT VARIANT 988 988  
FT VARIANT 989 989  
FT VARIANT 990 990  
FT VARIANT 991 991  
FT VARIANT 992 992  
FT VARIANT 993 993  
FT VARIANT 994 994  
FT VARIANT 995 995  
FT VARIANT 996 996  
FT VARIANT 997 997  
FT VARIANT 998 998  
FT VARIANT 999 999  
FT VARIANT 1000 1000  
FT VARIANT 1001 1001  
FT VARIANT 1002 1002  
FT VARIANT 1003 1003  
FT VARIANT 1004 1004  
FT VARIANT 1005 1005  
FT VARIANT 1006 1006  
FT VARIANT 1007 1007  
FT VARIANT 1008 1008  
FT VARIANT 1009 1009  
FT VARIANT 1010 1010  
FT VARIANT 1011 1011  
FT VARIANT 1012 1012  
FT VARIANT 1013 1013  
FT VARIANT 1014 1014  
FT VARIANT 1015 1015  
FT VARIANT 1016 1016  
FT VARIANT 1017 1017  
FT VARIANT 1018 1018  
FT VARIANT 1019 1019  
FT VARIANT 1020 1020  
FT VARIANT 1021 1021  
FT VARIANT 1022 1022  
FT VARIANT 1023 1023  
FT VARIANT 1024 1024  
FT VARIANT 1025 1025  
FT VARIANT 1026 1026  
FT VARIANT 1027 1027  
FT VARIANT 1028 1028  
FT VARIANT 1029 1029  
FT VARIANT 1030 1030  
FT VARIANT 1031 1031  
FT VARIANT 1032 1032  
FT VARIANT 1033 1033  
FT VARIANT 1034 1034  
FT VARIANT 1035 1035  
FT VARIANT 1036 1036  
FT VARIANT 1037 1037  
FT VARIANT 1038 1038  
FT VARIANT 1039 1039  
FT VARIANT 1040 1040  
FT VARIANT 1041 1041  
FT VARIANT 1042 1042  
FT VARIANT 1043 1043  
FT VARIANT 1044 1044  
FT VARIANT 1045 1045  
FT VARIANT 1046 1046  
FT VARIANT 1047 1047  
FT VARIANT 1048 1048  
FT VARIANT 1049 1049  
FT VARIANT 1050 1050  
FT VARIANT 1051 1051  
FT VARIANT 1052 1052  
FT VARIANT 1053 1053  
FT VARIANT 1054 1054  
FT VARIANT 1055 1055  
FT VARIANT 1056 1056  
FT VARIANT 1057 1057  
FT VARIANT 1058 1058  
FT VARIANT 1059 1059  
FT VARIANT 1060 1060  
FT VARIANT 1061 1061  
FT VARIANT 1062 1062  
FT VARIANT 1063 1063  
FT VARIANT 1064 1064  
FT VARIANT 1065 1065  
FT VARIANT 1066 1066  
FT VARIANT 1067 1067  
FT VARIANT 1068 1068  
FT VARIANT 1069 1069  
FT VARIANT 1070 1070  
FT VARIANT 1071 1071  
FT VARIANT 1072 1072  
FT VARIANT 1073 1073  
FT VARIANT 1074 1074  
FT VARIANT 1075 1075  
FT VARIANT 1076 1076  
FT VARIANT 1077 1077  
FT VARIANT 1078 1078  
FT VARIANT 1079 1079  
FT VARIANT 1080 1080  
FT VARIANT 1081 1081  
FT VARIANT 1082 1082  
FT VARIANT 1083 1083  
FT VARIANT 1084 1084  
FT VARIANT 1085 1085  
FT VARIANT 1086 1086  
FT VARIANT 1087 1087  
FT VARIANT 1088 1088  
FT VARIANT 1089 1089  
FT VARIANT 1090 1090  
FT VARIANT 1091 1091  
FT VARIANT 1092 1092  
FT VARIANT 1093 1093  
FT VARIANT 1094 1094  
FT VARIANT 1095 1095  
FT VARIANT 1096 1096  
FT VARIANT 1097 1097  
FT VARIANT 1098 1098  
FT VARIANT 1099 1099  
FT VARIANT 1100 1100  
FT VARIANT 1101 1101  
FT VARIANT 1102 1102  
FT VARIANT 1103 1103  
FT VARIANT 1104 1104  
FT VARIANT 1105 1105  
FT VARIANT 1106 1106  
FT VARIANT 1107 1107  
FT VARIANT 1108 1108  
FT VARIANT 1109 1109  
FT VARIANT 1110 1110  
FT VARIANT 1111 1111  
FT VARIANT 1112 1112  
FT VARIANT 1113 1113  
FT VARIANT 1114 1114  
FT VARIANT 1115 1115  
FT VARIANT 1116 1116  
FT VARIANT 1117 1117  
FT VARIANT 1118 1118  
FT VARIANT 1119 1119  
FT VARIANT 1120 1120  
FT VARIANT 1121 1121  
FT VARIANT 1122 1122  
FT VARIANT 1123 1123  
FT VARIANT 1124 1124  
FT VARIANT 1125 1125  
FT VARIANT 1126 1126  
FT VARIANT 1127 1127  
FT VARIANT 1128 1128  
FT VARIANT 1129 1129  
FT VARIANT 1130 1130  
FT VARIANT 1131 1131  
FT VARIANT 1132 1132  
FT VARIANT

DR InterPro: IPR007084; BRICHOS.  
 DR Pfam: PF04089; BRICHOS; 1.  
 DR PROSITE: PS50869; BRICHOS; 1.  
 KW Glycoprotein; Transmembrane; Cleavage on pair of basic residues;  
 Chondrogenesis.  
 CHAIN 1 210  
 FT PROPEP 211 214  
 FT CHAIN 215 333  
 FT TRANSMEM 42 62  
 FT DOMAIN 104 201  
 FT DISULFID 281 285  
 FT DISULFID 322 322  
 FT CARBOHYD 243 243  
 SQ SEQUENCE 333 AA; 36974 MW; 70B5FA9613DEAD CRC64;  
 N-LINKED (GLCNAC...) (POTENTIAL).  
 Query Match 28.2%; Score 492.5; DB 1; Length 333;  
 Best Local Similarity 33.4%; Pred. No. 5.3e-30;  
 Matches 110; Conservative 72; Mismatches 120; Indels 27; Gaps 12;  
 QY 6 PENCECHILNAFAFSKI-----CKSLKIGLVFGILATLIVLFW--GSKHFWPEVP 58  
 DB 15 PDVDFEC---GPPAYATVTVKPSGPAALLKGAVV--LISGAVLLFFGAIGAFYLMKSD 69  
 QY 59 KKAYDEHTFYSNGEKKIYMEIDPYTRTEIFRSNGTDELEVDHFNKGYGIYFVGLQ 118  
 DB 70 NHIYVHYMSINGKLQDSDMEIDANNLETFKMSGAEEALEVNDPFGNITGIRFAGE 129  
 QY 119 KCFIKQIV-IPEF-SEPEEED--ENEITTFPQSVIWEAKPIENRDLKNSK 173  
 DB 130 KCIYKQVAFVFEVETVQOSISELEKIMPVHEEALVWVAVGPPVDNSFL-SAR 188  
 QY 174 ILEICDNTVMYINPFLISVSELOPEEGEDLHPANEX--GIEONEQWVVCQAVE 230  
 DB 189 VLELCDDLEIFMLKPY--PKELQRRRVYAKVPTTKRHSPPRGNPG--PARWEND 244  
 QY 231 KTRHARQASEELPLINDYENG--IEFDMLEDERGYCCYCRGNRYCRVCEPLLYGP 288  
 DB 245 SRPSVQDESEPPNPDPYHGESEMTFDPRLDHEGICIECRSYTHCKICEPLGYNP 304  
 QY 289 YPYCYOGGVCIRVIMPCNMWVAPMLGRV 317  
 DB 305 WPIYNOGCRSACRVVWPCSMWVARILGVV 333  
 RESULT 10  
 CHM1 BRARE STANDARD; PRT; 286 AA.  
 ID CHM1 BRARE  
 AC P58239;  
 DT 16-OCT-2001 (Rel. 40, Created)  
 DT 16-OCT-2001 (Rel. 40, Last sequence update)  
 DT 15-MAR-2004 (Rel. 43, Last annotation update)  
 DE Chondromodulin-1 precursor (Chm-1) (Leukocyte cell-derived chemotaxin  
 1) (Contains: Chondrosulfactant protein (Ch-SP)).  
 DB 1) OR CHM1 OR CHM1.  
 OS Brachydanio rerio (Zebrafish) (Danio rerio).  
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 OC Actinopterygii; Neopterygii; Teleostei; Ostariophysi; Cypriniformes;  
 OC Cyprinidae; Danio.  
 OC NCBI\_TaxID=7955;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RX MEDLINE:21322690; PubMed:11429291;  
 RA Sachdev S.W., Dietz U.H., Oshima Y., Lang M.R., Knapik E.W.,  
 Hiraki Y., Shukunami C.;  
 "Sequence analysis of zebrafish chondromodulin-1 and expression  
 profile in the notochord and chondrogenic regions during cartilage  
 morphogenesis.";  
 Mech. Dev. 105:157-162(2001).  
 CC -1- FUNCTION: Bifunctional growth regulator. May contribute to the  
 rapid growth of cartilage and vascular invasion prior to the  
 replacement of cartilage by bone during endochondral bone  
 development (By similarity).

CC -1- SUBCELLULAR LOCATION: Cytoplasmic and secreted. Accumulated in the  
 CC inter-territorial matrix of cartilage (By similarity).  
 CC -1- PTM: After cleavage, the post-translationally modified Chm-1 is  
 CC secreted as a glycoprotein (By similarity).  
 CC -1- SIMILARITY: Belongs to the chondromodulin-1 family.  
 CC -1- SIMILARITY: Contains 1 BRICHOS domain.  
 CC -----  
 CC This SWISS-PROT entry is copyright. It is produced through a collaboration  
 CC between the Swiss Institute of Bioinformatics and the EMBL outstation  
 CC the European Bioinformatics Institute. There are no restrictions on its  
 CC use by non-profit institutions as long as its content is in no way  
 CC modified and this statement is not removed. Usage by and for commercial  
 CC entities requires a license agreement (See <http://www.isb-sib.ch/announce/>  
 CC or send an email to [license@isb-sib.ch](mailto:license@isb-sib.ch)).  
 CC -----  
 CC EMBL: AF322374; AK77023.1; -  
 CC ZFIN; ZDB-GENE-010713-1; lect1.  
 DR InterPro: IPR007084; BRICHOS.  
 DR Pfam: PF04089; BRICHOS; 1.  
 DR PROSITE: PS50869; BRICHOS; 1.  
 KW Glycoprotein; Transmembrane; Cleavage on pair of basic residues;  
 KW Chondrogenesis.  
 CHAIN 1 165  
 FT PROPEP 166 169  
 FT CHAIN 170 286  
 FT TRANSMEM 29 49  
 FT DOMAIN 75 157  
 FT DISULFID 234 238  
 FT DISULFID 235 235  
 FT CARBOHYD 191 191  
 SQ SEQUENCE 286 AA; 32235 MW; 21B377E034EEDB3 CRC64;  
 Query Match 23.1%; Score 403.5; DB 1; Length 286;  
 Best Local Similarity 32.3%; Pred. No. 2.2e-23;  
 Matches 108; Conservative 40; Mismatches 113; Indels 73; Gaps 14;  
 QY 7 ENCEDCHILNAFAFSKICKSLKIGLVFGILATLIVLFW--GSKHFWPEPKKAYLM 64  
 DB 3 ELAKTIPVEAVSLTKRGKLMKKKTLVAF--IAGALLFFGVGVAFYLMKTEKAISA 60  
 QY 65 EHTFYSNGEKKIYMEIDPYTRTEIFRSNGTDELEVDHFNKGYGIYFVGLQCFIKT 124  
 DB 61 H-----LGRNIDIKIENDS-----DSAGET--IVEVDFKAGITAVVFPGEKCFIKS 106  
 QY 125 QIKVIPFSEPEEIDENEITTFPQSVIWEAKPIENRDLKNSKILEICDNVTWY 184  
 DB 107 QART--ELSEDEAGV--KAEVA-----SLWITSEPLKDSFL--SPELLRCADLPYI 155  
 QY 185 WINPTLISVSELODFEEGEDLHPANEX-----KGIBQNBQWVVPQVK 228  
 DB 156 W-----HHFANSRALRKRSATMRQTSAGV--NRQPARRRS 192  
 QY 229 VEKTRHARQASEELPLINDYEN-GIE-----DPMMDERGYCCYCRGNRYCRVCEPL 283  
 DB 193 TASARBERPGEFYNENPHQNGSEGTWVPMLDHGHICCTECHRSTYTCERVCPL 252  
 QY 284 LGYYPYCYOGGVCIRVIMPCNMWVAPMLGRV 317  
 DB 253 GGYWPMVYVHGRPCRLIMPQWMAVARVLGVV 286  
 RESULT 11  
 SPB8 HUMAN STANDARD; PRT; 374 AA.  
 ID SPB8 HUMAN  
 AC P50452;  
 DT 01-OCT-1996 (Rel. 34, Created)  
 DT 01-OCT-1996 (Rel. 34, Last sequence update)  
 DT 28-FEB-2003 (Rel. 41, Last annotation update)  
 DE Cytoplasmic antipeptidase 2 (CAP2) (CAP-2) (Protease inhibitor 8)  
 DE (Serpin B8).  
 GN SERPINB8 OR P18.  
 OS Homo sapiens (Human).

```

OC Eukaryota; Metazoa; Chordata; Craniala; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Placenta;
RX MEDLINE=96102039; PubMed=8530382;
RA Sprecher C.A., Morgenstern K.A., Mathewes S., Dahlen J.R.,
RA Schrader S.K., Foster D.C., Kissel W., and partial characterization of two
RT "Molecular cloning, expression, and partial characterization of two
RT novel members of the ovalbumin family of serine proteinase
RT inhibitors.";
RL J. Biol. Chem. 270:29854-29861 (1995).
CC -1- SUBCELLULAR LOCATION: Cytoplasmic.
CC -1- SIMILARITY: Belongs to the serpin family. Ov-serpin subfamily.
CC -----
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL Outstation -
CC the European Bioinformatics Institute. There are no restrictions on its
CC use by non-profit institutions as long as its content is in no way
CC modified and this statement is not removed. Usage by and for commercial
CC entities requires a license agreement (see http://www.isb-sib.ch/announce/
CC or send an email to license@isb-sib.ch).
CC -----
DR EMBL; L40377; AAC41939.1; -.
DR PIR; A59273; A59273.
DR HSSP; P05120; 1BY7.
DR Genew; HGNC:8952; SERPINB8.
DR MIM; 601687;
DR GO; GO:0005829; C:cytosol; TAS.
DR GO; GO:0005515; F:protein binding; TAS.
DR GO; GO:0004868; F:serpin; TAS.
DR InterPro; IPR00215; Serpin.
DR Pfam; PF00079; serpin; 1.
DR SMART; SMO0093; SERPIN; 1.
DR PROSITE; PS00284; SERPIN; 1.
KW Serpin, Serine protease inhibitor.
FT ACT_SITE 339 340
SQ SEQUENCE 374 AA; 42786 MW; E855A03519AC60E CRC64;
Query Match 5.6%; Score 97.5; DB 1; Length 374;
Best Local Similarity 21.3%; Pred. No. 3,6;
Matches 63; Conservative 42; Mismatches 112; Indels 79; Gaps 14
QY 7 ENCEDC-HIINAEPKSKTKICKSLKIGLVGIALTLI--VLFWSGKHPVPEYKAY 62
DB 123 EDTCECRKHINPMVAEKTEGKSEVLDAIGVDPITKLVLVNAIYFKGK--WNBOFDKX 179
QY 63 DMEHFFYNGEKKIYMEIDPTREIFRSGSGTDETLVHDFKNGYGIYVGLQKPI 122
DB 180 TRGMLFKTNEEKIVQM-----MFEAK-----FKNGYAD-----EV 211
QY 123 KTIQKVIDEFSBDEEI-----DENBEITTT--FEQSVIWPAPKPIENF--DFL 169
DB 212 HTQVLELPV--FEELSWVILPDDNDFLAVVEKALTYEKKKATNSSEKTKSKYQVFL 268
QY 170 KNSKLTICDNTVMWNPITLSVSELQDFEEEGDLDHPAEKK-----GIQCN 219
DB 269 PRLKEE-----SYDLEPFLRLKGLDAFDPAKADFSQMSSTEKVPLSKVAHKCPVEZN 322
QY 220 EQMVPQVVEKTRARQASEBELPINDYTGEGIDFDPML-----DERGYCIYCR 271
DB 323 EGGTAAATAATAVNRSSRCRMERPQADH-----PLFPIRRHKTKCILFCGR 370
RESULT 12
ID YXDA_MYCCA STANDARD; PRT; 655 AA.
AC P45615;
DT 01-NOV-1995 (Rel. 32, Last sequence update)
DT 10-OCT-2003 (Rel. 42, Last annotation update)
DE Hypothetical lipoprotein in kdsB 5' region precursor (ORF).

```

```

OS      Mycoplasma capricolum.
CC      Bacteria; Firmicutes; Mollicutes; Mycoplasmataceae; Mycoplasma.
OK      NCBI_TaxID=2095;
RN      [1]
RP      SEQUENCE FROM N.A.
RC      STRAIN=ATCC 27343 / K1d;
RX      MEDLINE=95216643; PubMed=7703858;
RA      Zhu P.-P., Reizer J., Petekofsky A.;
RT      "Unique dicistronic operon (pstI-crr) in Mycoplasma capricolum
RT      encoding enzyme I and the glucose-specific enzyme IIA of the
RT      phosphoenolpyruvate:sugar phosphotransferase system: cloning,
RT      sequencing, promoter analysis, and protein characterization.";
RL      Protein Sci. 3:2115-2128(1994).
CC      -1- SUBCELLULAR LOCATION: Attached to the membrane by a lipid anchor
CC      (potential).
CC      -1- SIMILARITY: BELONGS TO THE MG067 / MG068 / MG395 FAMILY.
CC      -----
CC      This SWISS-PROT entry is copyright. It is produced through a collaboration
CC      between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC      the European Bioinformatics Institute. There are no restrictions on its
CC      use by non-profit institutions as long as its content is in no way
CC      modified and this statement is not removed. Usage by and for commercial
CC      entities requires a license agreement (See http://www.ebi.ac.uk/announcements/
CC      or send an email to license@ebi.ac.uk).
CC      -----
CC      EMBL; U15110; AAA70404.1; -.
CC      PIR; A57681; A57681.
CC      InterPro; IPR002414; DUF30/31.
CC      InterPro; IPR000437; Prok_LipoProt_S.
CC      Pfam; PF01732; DUF31.1
CC      PRINTS; PR00540; Y06768FAMILY.
CC      PROSITE; PS00013; PROKAR_LIPOPROTEIN_1.
CC      DR      Hypothetical protein; Lipoprotein; Membrane; signal; Palmitate.
CC      FT      SIGNAL
CC      FT      1 23
CC      FT      CHAIN
CC      FT      24 655
CC      FT      LIPID
CC      FT      24 24 N-palmitoyl cysteine (Potential).
CC      FT      LIPID
CC      FT      24 24 S-diacylglycerol cysteine (Potential).
CC      FT      SQ
CC      FT      SEQUENCE 655 AA; 75908 MW; A03E4B0EADB7B060 CnC64;
CC      -----
Query Match 5.6%; Score 97; Length 655;
Best Local Similarity 21.3%; Pred. No. 7.8;
Matches 57; Conservative 44; Mismatches 99; Indels 68; Gaps 12;

7 ENCEDCHILNAEAKRSKKICSLKICGLVFILTLTLVFWSGKHFMPEVPRKAYIMEH 66
155 DNIEDIIDSKLEILSKYI-----PQHNHPDYVK----- 185
67 TFVSGEKKIYMEIDPYTR-----EIPRSGNGIDETELVDFKNGYTGIFVG 116
186 --FKVSAEELIYKLYDRTFSIKFLVLYKDGGLSLNGSTGWLMDYHKYS--INKY-- 238
117 LQCFIKTQIVIPFSEPEEIDENIEITTFEEQ--YIWPDAEPIENRDPKLSK 173
239 --KFIATNLHLVLDNFNSLND--DQNKFF--NYYPDSSNKYIGGLGADNVTFSRKN 293
174 ILEIDQNTWMIWINDTLVSGLDQDFEEGEDLHPANE--KGIQEQNQWVVPQYKYEKT 232
294 NKSSENNIANYTLNN-----QDFEYLYLKNQDWSVYKFSKGLSE--PKYIFGAVDFPKD 344
233 R---HARQASEELPIINDYENGIEFD 256
345 RAIKHYEALQKEALINYNYKKNNEIN 372

RESULT 13
RRLP_VSVUO STANDARD; FRT; 2109 AA.
AC P16379;
DT 01-AUG-1990 (Rel. 15, Created)
DT 01-AUG-1990 (Rel. 15, Last sequence update)
DT 28-FEB-2003 (Rel. 41, Last annotation update)
DE RNA polymerase beta subunit (EC 2.7.7.46) (large structural protein)

```

```

DE (L protein).
GN L.
OS Vesicular stomatitis virus (serotype New Jersey / strain Ogden).
OC Viruses; ssRNA negative-strand viruses; Mononegavirales;
OC Rhabdoviridae; Vesiculoviruses.
OC NCBI_TaxID=11283;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=90177235; PubMed=2155516;
RA Barik S., Rud E.W., Luk D., Banerjee A.K., Kang C.Y.;
RA "Nucleotide sequence analysis of the L gene of vesicular stomatitis
RT virus (New Jersey serotype): identification of conserved domains in L
RL proteins of nonsegmented negative-strand RNA viruses."
RT Virology 175:332-337(1990).
CC -1- FUNCTION: THIS PROTEIN IS PROBABLY A COMPONENT OF THE ACTIVE
CC POLYMERASE. IT MAY FUNCTION IN RNA SYNTHESIS, CAPPING, AS WELL AS
CC METHYLATION OF CAPS, AND POLY(A) SYNTHESIS.
CC -1- CATALYTIC ACTIVITY: N nucleoside triphosphate = N diphosphate +
CC (RNA) (N).
CC -1- SUBUNIT: THOUGHT TO FORM A TRANSCRIPTION COMPLEX WITH THE
CC NUCLEOCAPSID (N) PROTEIN.
CC -1- SIMILARITY: WITH THE L PROTEIN OF OTHER RHABDOVIRUSES AND
CC PARAMYXOVIRUSES.
CC CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation-
CC the European Bioinformatics Institute. There are no restrictions on its
CC use by non-profit institutions as long as its content is in no way
CC modified and this statement is not removed. Usage by and for commercial
CC entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC or send an email to license@isb-sib.ch).
CC
CC EMBL: M29786; AAA48442.1; -.
DR PIR: A46309; A46309.
DR InterPro: IPR007098; RNA_pol_monon.
DR InterPro: IPR002877; RrmJ_Fcbl.
DR InterPro: IPR001016; Viral_RNA_pol_L.
DR Pfam: PF01728; Fted; 1.
DR Pfam: PF00946; Paramyx_RNA_pol; 1.
DR Transferrase; RNA-directed RNA polymerase.
KM SEQUENCE 2109 AA; 24211 MW; 724CF90BCE26CAB9 CRC64;
SQ
Query Match 5.5%; Score 96; DB 1; Length 2109;
Best local similarity 22.1%; Pred. No. 39;
Matches 52; Conservative 46; Mismatches 91; Indels 46; Gaps 13;
QY 55 PEVPKAYMEHTFFSYNGEKKIYMEIDPV-TTTFIRSGNGIDETLEVA-DPKA-GYTG 111
DB 306 PKIPKPPHEEHV---KGSVQELTORSNRIQTLIDIMSKVDLVLVYVGSFRHWGHP 362
QY 112 I-YFVGLOKCFIKTOIKVPEFSEPEEIDEN--EITTF-----FEOSVIVW--- 157
DB 363 IDVFOGLEK-LITQVNM-----EKHIDKEYPQLASDLARIVANKPSSSKWFPDP 413
QY 158 -----PAEKPIENDFLKNSKILEICNVVTWYINPTLISVSELQDFEESG---EDIH 207
DB 414 SKSPKHPFEYHVINKTWPAKIQDFGN---WKLPLIQCFEIPDLIDISVLYSDS 469
QY 208 FPAHEKKGE---QNEQWVVPQVKEKTHARQASHEELPINDYENGIEPDPM 259
DB 470 HSNMKKEVIOHVRSKRPNIPISNVKVLQTMLTNPAIMWKAFLKIDENGDDDDLI 524

```

```

OC Archaea; Euryarchaeota; Methanococci; Methanococcales;
OC Methanocaldococcaceae; Methanocaldococcus.
OC NCBI_TaxID=2190;
RN [1]
RP SEQUENCE FROM N.A.
RX STRAIN=JAL-1 / DSM 2661 / ATCC 43067;
RX MEDLINE=96337999; PubMed=8688087;
RA Bult C.J., White O., Olsen G.J., Zhou L., Fleischmann R.D.,
RA Sutton G.G., Blake O., Fitzgerald L.M., Clayton R.A., Gocayne J.D.,
RA Kerlavage G.A., R. Dougherty B.A., Tomb J.-F., Adams M.D., Reich C.I.,
RA Overbeek R., Kirkness E.F., Weinstock K.G., Merrick J.M., Glodek A.,
RA Scott J.L., Geoghegan N.S.M., Weidman J.F., Rhihmann J.L., Nguyen D.,
RA Utermarck T.R., Kelley J.M., Peterson J.D., Sedow P.W., Hanna M.C.,
RA Cotton M.D., Roberts K.W., Hurst M.A., Kaine B.P., Borodovsky M.,
RA Klenk H.-P., Fraser C.M., Smith H.O., Woese C.R., Venter J.C.;
RT "Complete genome sequence of the methanogenic archaeon, Methanococcus
RT jannaschii."
RT Science 273:1058-1073(1996).
CC -1- SIMILARITY: TO A.FULGIDUS AF0817.
CC
CC CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation-
CC the European Bioinformatics Institute. There are no restrictions on its
CC use by non-profit institutions as long as its content is in no way
CC modified and this statement is not removed. Usage by and for commercial
CC entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC or send an email to license@isb-sib.ch).
CC
CC EMBL: U67596; AAB95974.1; -.
DR PIR: A64494; A64494.
DR TIGR: MJ1554; -.
DR InterPro: IPR000437; Prok_Lipoprot_S.
KM Hypothetical protein; Signal; Complete proteome.
FT SIGNAL 1 20
FT CHAIN 21 622
SQ SEQUENCE 622 AA; 72282 MW; 86CAA62P3854D73F CRC64;
Query Match 5.5%; Score 95.5; DB 1; Length 622;
Best local similarity 22.0%; Pred. No. 9.4;
Matches 56; Conservative 40; Mismatches 79; Indels 79; Gaps 13;
QY 21 KSKTKCSLKIKGLVPSGIALTLVLFWGSKHEW---PEVPKAYMEHTFFSYNGEKK 76
DB 132 KYAKIKINISCGGLY-LTNMTLIVISMNKITSYVSNPMPKRIWMD---LNGS--- 183
QY 77 IYMEIDPVTRREIRSGNGT-----DETLEVDPFNGYTGIFY----- 114
DB 184 -YVD-----SRLYNGTLVLYVRKNSIDCPVWNNYKIGDKYIIPBLPYSMDF 233
QY 115 -----VGLQKCFIKTOIKVPEFSEPEEIDEN--EITTF-----FEOSVIVW--- 155
DB 234 TTYIIRINIKSGKVENSIIVGNYKTLVYKNNLFAVYMLKINKKMLNPLNBSADK 293
QY 156 WVPAA-----KPIENDFLKNSKILEICNVVTWYINPTLISVSELQDFEESG---EDIH 202
DB 294 YFPEVADKIKRVLENDFGNAKFEVETTERYL--SSLPSEKRNLMKKQLQNDPENY 351
QY 203 GEDLHFPANKEKGI 216
DB 352 LEE-HMEEFETGI 364

```

```

RESULT 14
YF54 METUA
ID YF54 METUA STANDARD; PRT; 622 AA.
AC OS8926;
DT 15-JUL-1998 (Rel. 36, Created)
DT 15-JUL-1998 (Rel. 36, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Hypothetical protein MJ1554 precursor.
GN MJ1554.
OS Methanococcus jannaschii.

```

```

RESULT 15
SPR2 CAEEL
ID SPR2 CAEEL STANDARD; PRT; 312 AA.
AC Q18240; O9PRJ0;
DT 10-OCT-2003 (Rel. 42, Created)
DT 10-OCT-2003 (Rel. 42, Last sequence update)
DT 10-OCT-2003 (Rel. 42, Last annotation update)
DE Suppressor of presenilin 2.
GN SPR-2 OR C2787.1.
OS Caenorhabditis elegans.
OC Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida; Rhabditidae;

```



CC Rhabdittidae; Feloderinae; Caenorhabditis.  
OX NCBI\_TaxID=6239;  
[1]  
RP SEQUENCE FROM N.A., FUNCTION, SUBCELLULAR LOCATION, AND TISSUE  
RP SPECIFICITY  
RC STRAIN=Bristol N2;  
RX MEDLINE=20570513; PubMed=1114162;  
RA Wen C., Levitan D., Li X., Greenwald I.;  
RT "egl-2, a suppressor of the egg-laying defect caused by loss of sel-12  
RT presentin in Caenorhabditis elegans, is a member of the SET protein  
RT subfamily.";  
RI Proc. Natl. Acad. Sci. U.S.A. 97:14524-14529(2000).  
[2]  
RP SEQUENCE FROM N.A.  
RC STRAIN=Bristol N2;  
RA Matthews P.;  
RI Submitted (SEP-1995) to the EMBL/GenBank/DBJ databases.  
[3]  
RP REVISIONS.  
RI Durbin R.;  
RI Submitted (JUN-2001) to the EMBL/GenBank/DBJ databases.  
CC -1- FUNCTION: Potential corepressor protein, which may affect  
CC chromatin structure and/or transcription. May participate in the  
CC transcriptional repression of the presentin protein hop-1.  
CC -1- SUBCELLULAR LOCATION: Nuclear.  
CC -1- TISSUE SPECIFICITY: Widely expressed.  
CC -1- MISCELLANEOUS: Loss of function results in a suppression of sel-12  
CC mutant phenotypes, possibly by upregulating hop-1 expression.  
CC -1- SIMILARITY: Belongs to the nucleosome assembly protein (NAP)  
CC family.  
CC  
CC This SWISS-PROT entry is copyright. It is produced through a collaboration  
CC between the Swiss Institute of Bioinformatics and the EMBL outstation  
CC the European Bioinformatics Institute. There are no restrictions on its  
CC use by non-profit institutions as long as its content is in no way  
CC modified and this statement is not removed. Usage by and for commercial  
CC entities requires a license agreement (See <http://www.isb-sib.ch/announce/>  
CC or send an email to [license@isb-sib.ch](mailto:license@isb-sib.ch)).  
CC  
CC EMBL; AF221546; AAG2102.1; -.  
DR EMBL; Z54236; CA90979.2; -.  
DR WormPep; C2787.1; CE27804.  
DR InterPro; IPR002164; NAP\_family.  
DR Pfam; PF00956; NAP; 1.  
KM Transcription regulation; Repressor; Nuclear protein.  
FT DOMAIN 214 300  
FT GLU-RICH.  
SQ SEQUENCE 312 AA; 35974 MW; 0CB087474841ADC CRC64;  
  
Query Match 5.4%; Score 95; DB 1; Length 312;  
Best Local Similarity 22.4%; Pred. No. 4.4;  
Matches 47; Conservative 38; Mismatches 55; Indels 70; Gaps 12;  
  
QY 100 LEVHDFNGGTGYFVGLQCFIKTKQKVPFSEPEEBIDENBITTFFEQS----- 153  
DB 109 LEVOEFED-----LRSGFKITMF-DPWEYF-TNEVITKSHLOSSEPTB 152  
QY 154 ---VIWVPAEKPIENRD-----FLK-----NSKILEICD-----NVTWYINPTL 190  
DB 153 ITTELEWENKPKPPPEPGDSAHFTLEWLYAALPDSDEIAEVTKDLYVNPLOYYWMP-- 210  
QY 191 ISVSELODFEEBGDLHFPANEKKGIHQEQW-----VVPQVVEKTRHA---RQASEEE 242  
DB 211 ---DMQVEEE--DDIEDFLNERGVDENGQRIPRNISDSLKYDQESADGEGEDEEE 264  
QY 243 LPINDYTNGI-----EPDMLDERG 263  
DB 265 EDWGEHEHDGVEEFAEGEHEEDGAIIEEG 294

Search completed: March 30, 2004, 14:31:33  
Job time : 19 secs

Result No.	Score	Query Match	Length	DB	ID	Description
1	1709	97.9	317	6	Q9SM14	Q9SM14 equus caball
2	741	42.5	135	11	Q8CE74	Q8CE74 mus musculus
3	545	39.2	338	13	O7ZG92	O7ZG92 xenopus laevis
4	521	31.2	29.9	11	Q86UX1	Q86UX1 mus musculus
5	518.5	29.7	333	4	O8TAY6	O8TAY6 homo sapien
6	264	15.1	60	6	Q9S1O7	Q9S1O7 equus caball
7	112.5	6.4	489	16	Q9S1J5	Q9S1J5 clostridium
8	108	6.2	545.5	5	Q8U459	Q8U459 plasmodium
9	108	6.2	5507	5	Q81FN3	Q81FN3 plasmodium
10	105.5	6.0	2227	5	Q81F21	Q81F21 plasmodium
11	104.5	6.0	4994	17	Q8F019	Q8F019 methanosarc
12	103.5	5.9	938	5	Q81H10	Q81H10 plasmodium
13	103	5.8	1248	5	Q81E04	Q81E04 plasmodium
14	101.5	5.8	1612	5	Q81FN0	Q81FN0 plasmodium
15	100.5	5.8	265	16	Q8SENF6	Q8SENF6 oceanobact
16	100.5	5.8	350	5	Q9UAY5	Q9UAY5 caenorhabdi

17	100	5.7	827	16	Q9GQV4	Q9GQV4 mycoplasma
18	100	5.7	844	10	O04498	O04498 arabidopsis
19	99	5.7	679	5	O8ITE0	O8ITE0 plasmodium
20	98.5	5.6	487	5	O8MY47	O8MY47 plasmodium
21	98.5	5.6	487	5	O8II16	O8II16 plasmodium
22	98	5.6	1253	5	O8IKA0	O8IKA0 plasmodium
23	98	5.6	2109	12	O8BS44	O8BS44 vesicular s
24	98	5.6	16215	5	Q9NFS3	Q9NFS3 drosophila
25	98	5.6	18074	5	Q9ITU4	Q9ITU4 drosophila
26	97.5	5.6	149	11	O62810	O62810 rattus norv
27	97.5	5.6	309	4	Q9RAK1	Q9RAK1 homo sapien
28	97.5	5.6	494	17	O8PXB4	O8PXB4 methanosarc
29	97.5	5.6	494	17	O8PUY6	O8PUY6 methanosarc
30	97.5	5.6	799	16	Q9PR12	Q9PR12 ureaplasma
31	97	5.6	164	4	O8KXP6	O8KXP6 homo sapien
32	97	5.6	3901	5	Q9NE53	Q9NE53 caenorhabdi
33	96.5	5.5	313	11	O3S531	O3S531 rattus norv
34	96.5	5.5	374	4	Q7Z2V6	Q7Z2V6 homo sapien
35	96.5	5.5	494	17	O8RTG5	O8RTG5 methanosarc
36	96.5	5.5	494	17	O8TRX0	O8TRX0 methanosarc
37	96.5	5.5	497	17	O8TMH9	O8TMH9 methanosarc
38	96.5	5.5	513	17	O8TS61	O8TS61 methanosarc
39	96.5	5.5	732	17	P9S869	P9S869 sulfolobus
40	96.5	5.5	752	16	Q9GCG9	Q9GCG9 lactococcus
41	95.5	5.5	837	5	Q2S751	Q2S751 plasmodium
42	95.5	5.5	1090	16	O9CFI3	O9CFI3 lactococcus
43	95.5	5.5	1218	5	O8IDC8	O8IDC8 plasmodium
44	95	5.5	844	10	P92948	P92948 arabidopsis
45	94.5	5.4	658	16	O2S252	O2S252 helicobacte

## ALIGNMENTS

## RESULT

ID	Q95M14	PRELIMINARY;	PRT;	317	AA
----	--------	--------------	------	-----	----

DT 01-DEC-2001 (TREMBlRel. 19, Created)

DT 01-OCT-2003 (TREMBLE1, 25, Last annotation update)

GN TMD.

OC Eukaryota; Metazoa; Choc

0X NCBI TaxID=9796;

RP	SEQUENCE FROM N.A.
RM	(11)

RC STRAIN=111010ugibz  
RA Hasegawa T.!

phenomodulin."! BT  
KI  
"Molecular cloning and characterization of human

DR EMBL: AB059407: BAB69482.2: - - -  
RL submitted (APR-2001) CO LINE EMBL/GenBank/DBCP caracascc  
SUBMITTED

GO; GO:0006118: P:electron transport; IEA;

DR InterPro: IPR007084; BRICHOS.  
DR InterPro: IPR000345; CytoC heme BS.

```

BR      P1am; PF04089; BR1CHOS; 1.
BR      PROSITE: PS00190: CYTOCHROME C; 1

```

SO SEQUENCE 317 AA; 3/165 MW; 0B/CA9D0FE1ZDA86 CRC84;

Query Match	Similarity	Score	DB	Length
97.9%	1709	6	317	
97.2%	355	132		

Matches	308;	conservative	4;	mismatches	5;	indels	0;	gaps	0;
---------	------	--------------	----	------------	----	--------	----	------	----

QY 1 MAKNPENCDCCHLNAEAFKSKKICKSKLRLCGLVFGILALTLVLFWGSNHFWEVFNK 80

Db 1 MAKNPENCECHILNABAFKSKKICRSLKICGLVFGILALTLVLFWGNGHFWPEIFCA 80

61 AYDMEHTFYSNGEKKIYMEIDPVRTEIFRSGNGIDEIJEVHDFKNGYIGIYFVGLQRC 120

RP 5EQJUNC FROM N.A.  
 RX STRAIN-PVB/N-3: TISSUE-Breast tumor;  
 RX MEDLINE-22398257, PubMed-12677932;  
 RA Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G., Schuller G.D.,  
 RA Klausner R.D., Collins F.S., Wagner U., Shennett C.M., Schuller G.D.,  
 RA Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,  
 RA Hopkins R.F., Jordan H., Moore T., Vaux S.I., Wang J., Haefl F.,  
 RA DiCicco G.L., Maunula K., Farmer A.A., Rubin G.M., Hong L.,  
 RA Scapleton M., Soares W.B., Bonaldi M.F., Casavant T.L., Scheetz T.E.,  
 RA Brownstein M.C., Ussid T.B., Toshyski S., Carrinini P., Prange C.J.,  
 RA Raha S.S., Lofgren A.N., Peters G.J., Malek U.J., Garret P.H.,  
 RA Bosnak S.A., Mcwan P.J., McKernan K.J., Aramson R.D., Mullaly S.J.,  
 RA Rickard S., Worley K.C., Hale S., Garcia A.M., Gay L.U., Huijy S.W.,  
 RA Villalon D.C., Muzny P.C., Sodergren E.J., Lu X., Gibbs R.A.,  
 RA Fahy J., Helton E., Kettelman M., Maadan A., Rodrigues S., Sanchez A.,  
 RA Whiting M., Maadan A., Young A.C., Shevchenko Y., Bouffard G.G.,  
 RA Blakeley R.W., Touchman J.W., Green E.D., Dickson M.C.,  
 RA Rodriguez A.C., Grimwood J., Schmutz U., Myers R.M., Butterfield Y.S.,  
 RA Krzywnski M.I., Skaska U., Smailus D.E., Scherch A., Schein J.E.,

RA Jones S.J., Marra M.A.;  
 RT "Generation and initial analysis of more than 15,000 full-length human  
 RT and mouse cDNA sequences";  
 RL Proc. Natl. Acad. Sci. U.S.A. 99:16699-16903(2002).  
 RN [2]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=FVB/N-3; TISSUE=Breast tumor;  
 RA Strausberg R.;  
 RL Submitted (JAN-2003) to the EMBL/GenBank/DBJ databases.  
 DR EMBL; BC045152; AA45152.1; -  
 DR InterPro; IPR007084; BRICHOS.  
 DR Pfam; PF04089; BRICHOS; 1  
 SQ SEQUENCE 334 AA; 37225 MW; BFAA0B224DC9944 CRC64;

Query Match 29.8%; Score 521; DB 11; Length 334;  
 Best Local Similarity 36.3%; Pred. No. 1.3e-34;  
 Matches 119; Conservative 66; Mismatches 119; Indels 24; Gaps 11;

QY 6 PENECDHIINAFAFNSKIKCKS-----LKICGLVFGILALTLIVLFW--GSKHFWPEV 57  
 DB 15 PEDVEFC---SPPAYTTVTVPFGSGPTRLKVGAVV--LISGAVLLLFALGAFYFWKGN 69  
 QY 58 PKAYDEHTFYSNGEKKKIYMEIDPYTRTEIFRSGNGTDETLVHDFFKNGYTGIVFGL 117  
 DB 70 DNHIYVHYMSINGKLQDGSMEIDANNLETFKMSGAEALVAVDFKNGITGIRFAGG 129  
 QY 118 QKCFIKTOIKV-IPEF-SEPEEIDENE--ITTFEQSIVVPAEKPIENRDLKNSK 173  
 DB 130 EKCIKQVAKARIPEVGTATVKQISELEKIMPANEENLLIWAVDQVXSSFL-SSK 188  
 QY 174 ILEICDNVWYMINPTLISVSELQDFEEGEDLHPANEKKGLEQNEQWVVPQVKTEKR 233  
 DB 189 ILELCGLPIFWLKP--WPKETQREREVNRNSASPTTRPESRGNAGPRLSNTR 246  
 QY 234 -HARQASEBELPINDYTE--NGIEPDMDERGYCCICYCRNGRYCRVCEPLGYYPY 289  
 DB 247 PNVQDDAEPPNPDPHYQGESWTFDPRLDHEGICICIECRRSYTHCQKICEPLGYYFM 306  
 QY 290 PYCYQGRVICRVIMPCNMVWVARMGRV 317  
 DB 307 PYNVQGRSACRVIMPCSMVVARILGMV 334

## RESULT 5

Q8TAY6 PRELIMINARY; PRT; 333 AA.  
 AC Q8TAY6;  
 DT 01-JUN-2002 (TREMBlrel. 21, Created)  
 DT 01-JUN-2002 (TREMBlrel. 21, Last sequence update)  
 DT 01-MAR-2003 (TREMBlrel. 23, Last annotation update)  
 DE Chondromodulin I.  
 OS Homo sapiens (Human).  
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 OC Mammalia; Eutheria; Primates; Catarrhini; Homiidae; Homo.  
 OX NCBI\_TaxID=9606;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC TISSUE=Brain;  
 RA Strausberg R.;  
 RL Submitted (MAR-2002) to the EMBL/GenBank/DBJ databases.  
 DR EMBL; BC025659; AA425659.1; -  
 DR InterPro; IPR007084; BRICHOS.  
 DR Pfam; PF04089; BRICHOS; 1  
 SQ SEQUENCE 333 AA; 36974 MW; 40A81C68ADG65FDC CRC64;

Query Match 29.7%; Score 518.5; DB 4; Length 333;  
 Best Local Similarity 35.9%; Pred. No. 2e-34;  
 Matches 119; Conservative 65; Mismatches 119; Indels 27; Gaps 12;

QY 6 PENECDHIINAFAFNSKIKCKS-----LKICGLVFGILALTLIVLFW--GSKHFWPEV 58  
 DB 15 PEDVEFC---SPPAYTTVTVPFGSGPTRLKVGAVV--LISGAVLLLFALGAFYFWKGS 69

QY 59 KKAIDMEHTFYSNGEKKKIYMEIDPYTRTEIFRSGNGTDETLVHDFFKNGYTGIVFGLQ 118  
 DB 70 SHIYVHYTMSINGKLQDGSMEIDANNLETFKMSGAEALVAVDFKNGITGIRFAGGE 129  
 QY 119 KCFIKTOIKV-IPEF-SEPEEID---ENEITTFEQSIVVPAEKPIENRDLKNSK 173  
 DB 130 KCIYKQVAKARIPEVGTATVKQISELEKIMPANEENLLIWAVDQVXSSFL-SSK 188  
 QY 174 ILEICDNVWYMINPTLISVSELQDFEEGEDLHPANEKK--GIEQNEQWVVPQVKTE 230  
 DB 189 VLELCGLPIFWLKPETY--PREIQREREVNRNSASPTTRKPHSGPSNPG--AGRLLNE 244  
 QY 231 KTRHARQASEBELPINDYTE--IEPDMDERGYCCICYCRNGRYCRVCEPLGYYP 288  
 DB 245 TRPSVQEDSQAFNPDPNPHYQGESWTFDPRLDHEGICICIECRRSYTHCQKICEPLGYTP 304  
 QY 289 PYCYQGRVICRVIMPCNMVWVARMGRV 317  
 DB 305 PYNVQGRSACRVIMPCSMVVARILGMV 333

## RESULT 6

Q95LQ7 PRELIMINARY; PRT; 50 AA.  
 AC Q95LQ7;  
 DT 01-DEC-2001 (TREMBlrel. 19, Created)  
 DT 01-DEC-2001 (TREMBlrel. 19, Last sequence update)  
 DT 01-OCT-2003 (TREMBlrel. 25, Last annotation update)  
 DE Tenomodulin (Fragment).  
 GN TMND.  
 OS Equus caballus (Horse).  
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 OC Mammalia; Eutheria; Perissodactyla; Equidae; Equus.  
 OX NCBI\_TaxID=9796;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=Thoroughbred;  
 RA Hasegawa T.;  
 RL "Molecular cloning of equine tenomodulin gene partial cds."  
 RT Submitted (SEP-2001) to the EMBL/GenBank/DBJ databases.  
 DR EMBL; AB071388; BAB64928.1; -  
 DR GO; GO:0005489; F:electron transporter activity; IEA.  
 DR GO; GO:006118; P:electron transport; IEA.  
 DR InterPro; IPR000345; Cytochrome b5.  
 DR PROSITE; PS00190; CYTOCHROME\_C\_1.  
 FT NON TER 50  
 SQ SEQUENCE 50 AA; 5479 MW; B56BA16A56FB5756 CRC64;

Query Match 15.1%; Score 264; DB 6; Length 50;  
 Best Local Similarity 100.0%; Pred. No. 1.5e-14;  
 Matches 49; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCECHILNAFAFNSKIKCKSLKICGLVFGILALTLIVLFWG 49  
 DB 1 MAKNPENCECHILNAFAFNSKIKCKSLKICGLVFGILALTLIVLFWG 49

## RESULT 7

Q97LJ5 PRELIMINARY; PRT; 499 AA.  
 AC Q97LJ5;  
 DT 01-OCT-2001 (TREMBlrel. 18, Created)  
 DT 01-OCT-2001 (TREMBlrel. 18, Last sequence update)  
 DT 01-OCT-2003 (TREMBlrel. 25, Last annotation update)  
 DE Histidine kinase (HAMP, HiskA and Hapase domains).  
 GN CAC0565.  
 OS Clostridium acetobutylicum.  
 OC Bacteria; Firmicutes; Clostridia; Clostridiales; Clostridiaceae;  
 OC Clostridium.  
 OX NCBI\_TaxID=1488;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=ATCC 824 / DSM 792 / VKM B-1787;

RX MEDLINE=21359325; PubMed=1146286;  
 RA Noelling J., Breton G., Omeichenko M.V., Makarova K.S., Zeng Q.,  
 RA Gibson R., Lee H.M., Dubois J., Olu D., Hitti Y.I., Wolf Y.I.,  
 RA Tatusov R.L., Sabathe F., Doucelte-Stamm L., Soucaille P., Daly M.J.,  
 RA Bennett G.N., Koonin E.V., Smith D.R.,  
 RT "Genome sequence and comparative analysis of the solvent-producing  
 RT bacterium *Clostridium acetobutylicum*,"  
 RL J. Bacteriol. 183:4823-4838(2001).  
 DR EMBL; AE007571; AAK78544.1; --  
 DR PIR; E96969; E96969.  
 DR GO; GO:0016020; C:membrane; IEA.  
 DR GO; GO:0005524; F:ATP binding; IEA.  
 DR GO; GO:0016301; F:kinase activity; IEA.  
 DR GO; GO:0000155; F:two-component sensor molecule activity; IEA.  
 DR GO; GO:0000160; P:two-component signal transduction system (p. . .); IEA.  
 DR InterPro; IPR003594; ATPbind\_ATPase.  
 DR InterPro; IPR004358; Bact\_sens\_pr\_C.  
 DR InterPro; IPR003660; HAMF.  
 DR InterPro; IPR005467; His\_kinase.  
 DR InterPro; IPR003661; His\_kinase.  
 DR InterPro; IPR008358; Lantibiot\_regn.  
 DR Pfam; PF02518; HAMFase\_c\_1.  
 DR Pfam; PF00512; HisKA\_1.  
 DR PRINTS; PR00344; BCTRLSENSOR.  
 DR PRINTS; PR01780; LANTIBIOPROT.  
 DR SMART; SM00304; HAMF\_1.  
 DR SMART; SM00387; HAMFase\_c\_1.  
 DR SMART; SM00388; HisKA\_1.  
 DR PROSITE; PS00885; HAMF\_1.  
 DR PROSITE; PS0109; HIS\_KIN; 1.  
 DR KINASE; Complete proteome.  
 SQ SEQUENCE 499 AA; 57549 MW; 7FCA501BBD04F628 CRC64;

Query Match 6.4%; Score 112.5; DB 16; Length 499;  
 Best Local Similarity 22.3%; Pred. No. 0.71;  
 Matches 71; Conservative 49; Mismatches 103; Indels 95; Gaps 21;

QY 15 LNAAFAKSKKICKSKIKI-CGLVFGILALTLVL-----FMGSKFMEVP----- 58  
 DB 9 LKLYSRSKKKVKVSKIRLELVTFGICLLAFILGSGNTGYNDKMMABVDYSGIQGIS 68  
 QY 59 -----KKAVDMEHTFYSNGEKK-KIYNEIDPVTRTE--IFRSGNGDETLE 101  
 DB 69 SPMEDLKGMLDGKNSDSDEINKMIDGAKDIDKVL-----TDEGKVLFKSNNAEKKID 123  
 QY 102 VHD-----FKNGTGIYFVGLQKCFITKQIKVPERSEPEEID---ENE---ITT 147  
 DB 124 IHDLYKSKKFKREYDKATVINTVNGIIRKITSL--ESVEQYNEIDGKIKINKKAYITVS 181  
 QY 148 TFEQSVIIV-PAEKPIENRDEPLKNSKILICDNTWTW-----INPTLISVS 194  
 DB 182 GIPKQKITVIRGYSF-----FL--SGILAVITFFIFELPNRKMEYIESGSLVAIS 234  
 QY 195 ELQ-DPE--EEGEDLHPANEKKGKEQEQWVP---QYKVEKTRARQASEELPINDYT 249  
 DB 235 KNTLYRIMLWGED-----ELKKNLADINNSWASLEOKRIEKERAEKTK-----NDLI 282  
 QY 250 ENGIEED---PMLDERGY 264  
 DB 283 TN-VSHDLRTPLTSVKGY 299  
 RESULT 8  
 Q9U459 PRELIMINARY; PRT; 5458 AA.  
 AC Q9U459; Q9UPG4;  
 DT 01-MAY-2000 (TREMblrel. 13, Created)  
 DT 01-JUN-2001 (TREMblrel. 17, Last sequence update)  
 DT 01-OCT-2003 (TREMblrel. 23, Last annotation update)  
 DE Erythrocyte membrane-associated giant protein antigen 332.  
 OS Plasmodium falciparum.  
 OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.

OX NCBI\_Taxid=5833;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=FCCL/HN;  
 RA Shan Z.X., Yu X.B., Li X.R., Ma C.L., Fang J.M., Lu J.H., X J.;  
 RT "Molecular cloning and structure analysis of the Plasmodium falciparum  
 RT erythrocyte membrane-associated giant protein Ag332 (PF332) gene,"  
 RL Submitted (FEB-2001) to the EMBL/GenBank/DBJ databases.  
 DR EMBL; AF202180; AAF15293.3; --  
 DR DR GO; GO:0003723; F:RNA binding; IEA.  
 DR InterPro; IPR006763; Ag332.  
 DR InterPro; IPR001313; Pumilio/Puf.  
 DR Pfam; PF04671; Ag332; 179.  
 SQ SEQUENCE 5458 AA; 615269 MW; 4FEA37D92BD80172 CRC64;

Query Match 6.2%; Score 108; DB 5; Length 5458;  
 Best Local Similarity 24.4%; Pred. No. 33;  
 Matches 38; Conservative 37; Mismatches 47; Indels 34; Gaps 7;

QY 136 EBEIDENEITTTFFQSVI-----WVPAEKPIENRDFLNSKILICDNT----- 182  
 DB 3003 EQDVSDNEIIVE--ERSVIEAEENWVI--EKVEE-EEGLDNEVYDEEDSVSEQAEER 3056  
 QY 183 MWMINPTLISVSLQDPEEGEDLHPANE--KKGKEQNE-----QWVPOV 227  
 DB 3057 VLINEELKSSQVEDKVENELMNEBVEETQSVAENNEDEKELNDYVEETSVTEEV 3116  
 QY 228 KVEKTRARQASEELPINDYTNENGIEFDPMLDERG 263  
 DB 3117 VVDEVPNSKEVGEIESIEIEIIEVEDGLTDDLVGQCG 3152  
 RESULT 9

Q8IHN3 PRELIMINARY; PRT; 5507 AA.  
 ID Q8IHN3  
 AC Q8IHN3;  
 DT 01-MAR-2003 (TREMblrel. 23, Created)  
 DT 01-MAR-2003 (TREMblrel. 23, Last sequence update)  
 DT 01-JUN-2003 (TREMblrel. 24, Last annotation update)  
 DE Antigen 332, putative.  
 GN PF1\_0507.  
 OS Plasmodium falciparum (isolate 3D7).  
 OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.  
 CX NCBI\_Taxid=36329;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=3D7;  
 RK MEDLINE=22255705; PubMed=1236864;  
 RA Gardner M.J., Hall N., Fung E., White O., Berziman M., Hyman R.W.,  
 RA Carlson J.M., Pain A., Nelson K.E., Bowman S., Paulsen I.T., James K.,  
 RA Elesen J.A., Rutherford K., Salzberg S.L., Craig A., Kyes S.,  
 RA Chan M.-S., Nene V., Shallom S.J., Sub B., Peterson J., Angiuoli S.,  
 RA Pettes M., Allen J., Shellenberg J., Haft D., Mather M.W., Valdivia A.B.,  
 RA Martin D.M.A., Fairland A.H., Fraunholz M.J., Roos D.S., Ralph S.A.,  
 RA McRadden G.I., Cummings L.M., Subramanian G.M., Cunningham C.,  
 RA Venter J.C., Carucci P.J., Hoffman S.L., Newbold C., Davis R.W.,  
 RA Fraser C.M., Barrett B.;  
 RT "Genome sequence of the human malaria parasite Plasmodium  
 RT falciparum,"  
 RL Nature 419:498-511(2002).  
 DR EMBL; AB014843; AAN36076.1; --  
 DR InterPro; IPR006763; Ag332.  
 DR Pfam; PF04671; Ag332; 185.  
 SQ SEQUENCE 5507 AA; 621156 MW; 03B095165D1490AE CRC64;

Query Match 6.2%; Score 108; DB 5; Length 5507;  
 Best Local Similarity 24.4%; Pred. No. 33;  
 Matches 38; Conservative 37; Mismatches 47; Indels 34; Gaps 7;  
 QY 136 EBEIDENEITTTFFQSVI-----WVPAEKPIENRDFLNSKILICDNT----- 182  
 DB 2938 EQDVSDNEIIVE--ERSVIEAEENWVI--EKVEE-EEGLDNEVYDEEDSVSEQAEER 2991

```

QY 183 MYWNPFLISVSELODFEEGEDLHFPANE--KKGIEONE-----QWVPOV 227
DB 2992 VYINBELIKSSVDEVDKVENELMNEVNEFQSVANNEBDEKLDNYVEETSEV 3051
QY 228 KVEKTRHARQASEEELPINDYTENGIEFDPMLDERG 263
DB 3052 VVDEVNSKEVQIEIEIIEVEDGLTDLVGQCG 3087

RESULT 10
Q81121 ID Q81121 PRELIMINARY; PRT; 2227 AA.
AC Q81121;
DT 01-MAR-2003 (TREMBlrel. 23, Created)
DT 01-MAR-2003 (TREMBlrel. 23, Last sequence update)
DE Hypothetical protein.
GN Pf11.0354.
OS Plasmodium falciparum (isolate 3D7).
OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.
OX NCBI_TaxID=36329;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=3D7;
RX MEDLINE=22255705; PubMed=12368664;
RA Gardner M.J., Hall N., Fung B., White O., Berriman M., Hyman R.W.,
RA Carlton J.M., Pain A., Nelson K.E., Bowman S., Paulsen I.T., James K.,
RA Eisen J.A., Rutherford K., Salzberg S.L., Craig A., Kyes S.,
RA Chan M.-S., Nene V., Shallow S.J., Suh B., Peterson J., Angiuoli S.,
RA Partea M., Allen V., Selengut J., Haft D., Mather M.W., Valdivia A.B.,
RA Martin D.M.A., Fairlamb A.H., Fraunholz M.J., Roos D.S., Ralph S.A.,
RA McFadden G.I., Cummings L.M., Subramanian G.M., Mungall C.,
RA Venter J.C., Carucci D.J., Hoffman S.L., Newbold C., Davis R.W.,
RA Fraser C.M., Batteil B.;
RT "Genome sequence of the human malaria parasite Plasmodium
RL Nature 419:498-511 (2002).
RW EMBL: AE014841; AAM35938.1;
KW Hypothetical protein.
SQ SEQUENCE 2227 AA; 262840 MW; 6D5D4BEEF64339C CRC64;

Query Match 6.0%; Score 105.5; DB 5; Length 2227;
Best Local Similarity 18.3%; Pred. No. 17;
Matches 44; Conservative 56; Mismatches 64; Indels 77; Gaps 11;

QY 69 YSNGEKKTME-----IDPV--TRFE---IFRSGNGIDETLEVHDK 106
DB 1147 FSNMNDNKNVYMEGLINQDNVNNNNKSCDNIKMRKSLNFRSRSYGEHKSLDVY--- 1203
QY 107 NGYTGIVFGLQKCFITQIKVIEFSEPEE-----IDENEITTTFFEOGVIWVPAE 160
DB 1204 -----QCYVYVGN--KLINKVNDKKEEDNNNSYLNEDNNAQMFEEH-----NS 1246
QY 161 KP--TENRDLKSKIKIICDNTWMIWNPFLISVSELODFEEGEDLHFPANE 212
DB 1247 NPYVVDENNMKNVNNVNLVNNNSNYVDKSK-ENAEKSKSDILNENNIHILKQ 1305
QY 213 KKGIEQOWVPOVQKVEKTRHARQ-----ASEELPINDYTENG 252
DB 1306 KKKIQQNNEFISQADIENTRNSQEEVYKEHEPLWVINSNEKKSVEELIISDMSNR 1365
QY 253 I 253
DB 1366 V 1366

RESULT 11
Q8PU19 ID Q8PU19 PRELIMINARY; PRT; 494 AA.
AC Q8PU19;
DT 01-OCT-2002 (TREMBlrel. 22, Created)
DT 01-OCT-2002 (TREMBlrel. 22, Last sequence update)
DT 01-OCT-2002 (TREMBlrel. 22, Last annotation update)

```

```

DE Transposase.
GN MM2345.
OS Methanosarcina mazei (Methanosarcina frisia).
OC Archaea; Euryarchaeota; Euryarchaeota orders incertae sedis;
OC Methanosarcinales; Methanosarcinaceae; Methanosarcina.
OX NCBI_TaxID=2209;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=Goel / G1 / ATCC BAA-199 / DSM 3647 / OCM 88;
RX MEDLINE=22120827; PubMed=12125824;
RA Deppemeister U., Johann A., Hartesch T., Merl R., Schults R.A.,
RA Martinez-Ariza R., Heme A., Wietzer A., Baumeister S., Jacobi C.,
RA Bhuttagemah H., Lienard T., Christmann A., Boemcke W., Steckel S.,
RA Fritz H.-J., Gottschalk G.;
RT "The genome of Methanosarcina mazei: evidence for lateral gene
RT transfer between Bacteria and Archaea."
RL J. Mol. Microbiol. Biotechnol. 4:453-461 (2002).
DR EMBL: AE013477; AAM32041.1;
DR InterPro; IPR002559; Transposase_11.
DR Pfam; PF01609; Transposase_11; I.
KW Complete proteome.
SQ SEQUENCE 494 AA; 57328 MW; 0C6514323B167A5F CRC64;

Query Match 6.0%; Score 104.5; DB 17; Length 494;
Best Local Similarity 22.6%; Pred. No. 3.2; Indels 101; Gaps 14;
Matches 62; Conservative 30; Mismatches 81;

QY 56 EYFKKAYMEHTF--YSNGEKKTMEIDPVRTETFRSGNGIDETLEVHDKQGYTGY 113
DB 184 QIPFELVKKRKLKIKAKK-----LD-----EKKKQVITNDARIMYKD----- 228
QY 114 FVGLQKCIKQIKIPIFSEPEEIDENEITTTFFEOGVIWVPAEKINRDLKSK 173
DB 229 --GSKKPEYNCQAV-----DEKEQIT-----VAADVNEENDLHVEP 265
QY 174 ILEICDNTWMIWNPFLI-----SVSELODFEEGEDLHFPAN-----EKKGIEONEQW 222
DB 266 IIQNVKNTLGY--KPTIYLADAGFYSYENLEFLFEGSDAIPNNYAKREG----- 316
QY 223 VVPOVQKVEKTRHARQASEEELPINDYTENGIEFDPMLDERGYCCICYCRG--NRCCR 277
DB 317 -----KTRKFRKSL-----FKYSEKQ-----CYCPAFEFIPSRLOK 350
QY 278 RYCEPLIGYPPYPCYQ-----GGRVYCR 301
DB 351 RKGEPLDLYVCSYCSQCVLNACTESGKRTTR 384

RESULT 12
Q8IHVO ID Q8IHVO PRELIMINARY; PRT; 938 AA.
AC Q8IHVO;
DT 01-MAR-2003 (TREMBlrel. 23, Created)
DT 01-MAR-2003 (TREMBlrel. 23, Last sequence update)
DT 01-OCT-2003 (TREMBlrel. 25, Last annotation update)
DE Protein phosphatase 2C.
GN Pf11.0396.
OS Plasmodium falciparum (isolate 3D7).
OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.
OX NCBI_TaxID=36329;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=3D7;
RX MEDLINE=22255705; PubMed=12368664;
RA Gardner M.J., Hall N., Fung B., White O., Berriman M., Hyman R.W.,
RA Carlton J.M., Pain A., Nelson K.E., Bowman S., Paulsen I.T., James K.,
RA Eisen J.A., Rutherford K., Salzberg S.L., Craig A., Kyes S.,
RA Chan M.-S., Nene V., Shallow S.J., Suh B., Peterson J., Angiuoli S.,
RA Partea M., Allen V., Selengut J., Haft D., Mather M.W., Valdivia A.B.,
RA Martin D.M.A., Fairlamb A.H., Fraunholz M.J., Roos D.S., Ralph S.A.,
RA McFadden G.I., Cummings L.M., Subramanian G.M., Mungall C.,
RA Venter J.C., Carucci D.J., Hoffman S.L., Newbold C., Davis R.W.,

```

RA Frazer C.M., Barrell B.;  
 RT "Genome sequence of the human malaria parasite Plasmodium  
 falciptarum";  
 RL Nature 419:498-511 (2002).  
 DR EMBL: AE014842; AN35979.1; -  
 DR GO; GO:0008287; C:protein serine/threonine phosphatase complex; IEA.  
 DR GO; GO:0003824; C:catalytic activity; IEA.  
 DR GO; GO:0004722; F:protein serine/threonine phosphatase activity; IEA.  
 DR GO; GO:0006470; P:protein amino acid dephosphorylation; IEA.  
 DR InterPro: IPR000222; P2C.  
 DR InterPro: IPR001932; P2C-like.  
 DR Pfam: PF00481; P2C; 3.  
 DR SMART; SM00332; P2C; 1.  
 DR PROSITE; PS01032; P2C; 1.  
 SQ SEQUENCE 938 AA; 107250 MW; 75B1B743B5BA32 CRC64;

Query Match 5.9%; Score 103.5; DB 5; Length 938;  
 Best Local Similarity 26.2%; Pred. No. 8.6; Mismatches 67; Conservative 32; Indels 75; Gaps 16;

QY 33 GLVFGIALTLIVLFWGSKHFM--PEVPRKAYDMEHTYSNGEKKIYMEIDPVTRTE-- 88  
 DB 702 GRVDGNLNLTRA1---GDLHYKRDPLPQK-----DQKISAPPEITCVLLPED 747  
 QY 89 --IPRSGNGTDETLVHDFKNGVTGIYFVGLQKCFIKTQIKVPEF--SEPEEI----- 139  
 DB 748 EFLFLACD-----IWCKDGQD---VVG---FKXTRLEKFEIIPDPNNSLDNTENS 794  
 QY 140 -----DENEETITTFEFGSVIWPAAEKPI--ENRDFLNSKIT--EICDVTWYMI 186  
 DB 795 EHMNTNNNSDQNEKIKSE--DQNNINNEQENILNTSNDPQNEKYLNNENNNDINM--- 849  
 QY 167 NPTLISVSELQDFEEGEDLHFPANEKKGIEQNEQWVVPQVKVETRRARQASEE---EL 243  
 DB 850 -----QKSEETEDLIERDEK--HDAPVIERKKYKFKNLQICELCDEEC 896

QY 244 PINDYTEN-GIEFDPM 258  
 DB 897 LSNNKENDGICDMM 912

RESULT 13  
 Q81204 PRELIMINARY; PRT; 1248 AA.  
 AC Q81204;  
 DT 01-MAR-2003 (TrEMBLrel. 23, Created)  
 DT 01-MAR-2003 (TrEMBLrel. 23, Last sequence update)  
 DT 01-MAR-2003 (TrEMBLrel. 23, Last annotation update)  
 DE Hypothetical protein.  
 GN PF0100C.  
 OS Plasmodium falciptarum (isolate 3D7).  
 OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.  
 RX NCB1\_Taxid=36329;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RA MEDLINE:2225708; PubMed:1236867;  
 RA Hall N., Pain A., Berriman M., Churcher C., Harris B., Harris D.,  
 RA Mungall K., Bowman S., Atkin R., Baker S., Barron A., Brooks K.,  
 RA Buckee C.O., Burrows C., Cherevach I., Chillingworth C.,  
 RA Chillingworth T., Christodoulou Z., Clark R., Clark R., Corcoran C.,  
 RA Cronin A., Davies R., Davis P., Dear P., Dearden F., Doggett J.,  
 RA Fretwell T., Goble A., Goodhead I., Gilliam R., Hamlin N., Hance Z.,  
 RA Harper D., Hauser H., Hornsby T., Holroyd S., Horrocks P.,  
 RA Humphrey S., Jagels K., James K.D., Johnson D., Kechrou A.,  
 RA Knights A., Konfortov B., Kyes K.D., Larke N., Lawson D., Lennard N.,  
 RA Line A., Maddison M., McLean J., Mooney P., Moule S., Murphy L.,  
 RA Oliver K., Ormond D., Price C., Quail M.A., Rabinowitsch E.,  
 RA Rajandream M.A., Rutter S., Rutherford K.M., Sanders M., Simmonds M.,  
 RA Seeger K., Sharp S., Smith R., Squares R., Squares S., Stevens K.,  
 RA Taylor K., Tivey A., Unwin L., Whitehead S., Woodward J.,  
 RA Sulston J.E., Craig A., Newbold C., Barrell B.G;  
 RT "Sequence of Plasmodium falciptarum chromosomes 1, 3-9 and 13";  
 RL Nature 419:527-531 (2002).

DR EMBL: AL034557; CAD49112.1; -  
 KM Hypothetical protein.  
 SQ SEQUENCE 1248 AA; 154216 MW; A6EDA9493C71ACS CRC64;

Query Match 5.9%; Score 103; DB 5; Length 1248;  
 Best Local Similarity 22.2%; Pred. No. 13; Mismatches 74; Conservative 43; Indels 118; Gaps 17;

QY 7 ENCEDCHI-LNMAEF-----KSKIKCKSLKIGLVLATLTLVFWGSKHF 53  
 DB 895 EECKEEVQLNKLFLDSCIKELIKEKSKGCKMLB-----IYLDLKDRAV 941  
 QY 54 W-----PEVEKK-----AYDMEHTYSNGEKKIY-----N 79  
 DB 942 LINKLEMMKRIQKNSNNLFENIKTQVKNNENNYIEMMLNDNMKGMSENDCIFDEM 1001  
 QY 80 EIDPVTRTEIFRSGNGTDETLVHDFKNGVTGIYFVGLQKCFIKTQIKV----- 128  
 DB 1002 LNNMKSD--RIGQCKDNTIE-DNLKSNMNLGQVN-EKCLMEKINSNHWENIKN 1057  
 QY 129 IPEFSEPEEIDENEI-----TTTFE--QSVIWPAAEK-----PIENRDFLNS 172  
 DB 1058 CTGDEKEKEKIKVEMENIENIGSTGKIENIENIDEIVEEKYSFNVPEEENDEEKS 1117  
 QY 173 KILIECDNMTWVWNPITLISVSELQDFEEGEDLHFPANEKKGIEQNEQ-WVVPQVKVEK 231  
 DB 1118 KILNRSNIQSKLISLIDN-----EKGID-----EKSDNGNEIHW----- 1155

QY 232 TRHARQASEEELPINDYTENGIEFDPMLDERGY 264  
 DB 1156 ---NGLTLEIYKNSYTNISIEYIPIDEEY 1184

RESULT 14  
 Q81FNO PRELIMINARY; PRT; 1612 AA.  
 AC Q81FNO;  
 DT 01-MAR-2003 (TrEMBLrel. 23, Created)  
 DT 01-MAR-2003 (TrEMBLrel. 23, Last sequence update)  
 DT 01-MAR-2003 (TrEMBLrel. 23, Last annotation update)  
 DE Hypothetical protein.  
 GN PF0115C.  
 OS Plasmodium falciptarum (isolate 3D7).  
 OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.  
 RX NCB1\_Taxid=36329;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RA Hamlin N., Pain A., Berriman B., Hall N., Bowman S., Churcher C.,  
 RA Harris B., Harris D., Lawson D., Quail M., Barrell B.,  
 RA Submitted (FEB-1999) to the EMBL/Genbank/DBJ databases.  
 RL EMBL: AL035477; CAD49270.1; -  
 KW Hypothetical protein.  
 SQ SEQUENCE 1612 AA; 190946 MW; F004408BE4521D07 CRC64;

Query Match 5.8%; Score 101.5; DB 5; Length 1612;  
 Best Local Similarity 24.1%; Pred. No. 25; Mismatches 42; Conservative 35; Indels 70; Gaps 5;

QY 74 KKKIYMEIDPVTRTEIFRSGNGTDETLVHDFKNGVTGIYFVGLQKCFIKTQIKVPEFS 133  
 DB 1461 KKKINIDNDPINEI-----ENEKSKNNPVKFKFLSTLFI--LKYIDLVT 1503  
 QY 134 EPEEIDENEIITTFEFGSVIWPAAEKPIENRDFLNSKITLIECDNVTM-WVNPITLIS 192  
 DB 1504 ----LVKNNEFTDCDFLNTLMTQWDKVEKKSLNKGAVENDDINVEHINNEDEK 1558

QY 193 VSELQDFEEGEDLHFPANEKKGIEQNEQWVVPQVKVETRRARQASEELPIN 246  
 DB 1559 NEKLNKKGVEYDVTENLNQBAEEAEE---EAESEEEEDKFLTPHLPIN 1608

RESULT 15  
 Q8ENPF





GenCore version 5.1.6  
Copyright (c) 1993 - 2004 CompuGen Ltd.

OM protein - protein search, using sw model

Run on: March 30, 2004, 14:32:38 ; Search time 45 Seconds  
(without alignments)  
1843.262 Million cell updates/sec

Title: US-10-089-600-2  
Perfect score: 1745  
Sequence: 1 MARKPENECEDCHILNAEAF.....VICRVMPCNWMVARMIGRV 317

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 1065169 seqs, 261661801 residues

Total number of hits satisfying chosen parameters: 1065169

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%  
Listing first 45 summaries

Database :

Published Applications\_AA:\*

- 1: /cgn2\_6/ptodata/2/pubpaa/US07\_PUBCOMB.pep:\*
- 2: /cgn2\_6/ptodata/2/pubpaa/PCF\_NEW\_PUB.pep:\*
- 3: /cgn2\_6/ptodata/2/pubpaa/US06\_NEW\_PUB.pep:\*
- 4: /cgn2\_6/ptodata/2/pubpaa/US06\_PUBCOMB.pep:\*
- 5: /cgn2\_6/ptodata/2/pubpaa/US07\_NEW\_PUB.pep:\*
- 6: /cgn2\_6/ptodata/2/pubpaa/PCFUS\_PUBCOMB.pep:\*
- 7: /cgn2\_6/ptodata/2/pubpaa/US08\_NEW\_PUB.pep:\*
- 8: /cgn2\_6/ptodata/2/pubpaa/US08\_PUBCOMB.pep:\*
- 9: /cgn2\_6/ptodata/2/pubpaa/US09A\_PUBCOMB.pep:\*
- 10: /cgn2\_6/ptodata/2/pubpaa/US09B\_PUBCOMB.pep:\*
- 11: /cgn2\_6/ptodata/2/pubpaa/US09C\_PUBCOMB.pep:\*
- 12: /cgn2\_6/ptodata/2/pubpaa/US10A\_PUBCOMB.pep:\*
- 13: /cgn2\_6/ptodata/2/pubpaa/US10B\_PUBCOMB.pep:\*
- 14: /cgn2\_6/ptodata/2/pubpaa/US10C\_PUBCOMB.pep:\*
- 15: /cgn2\_6/ptodata/2/pubpaa/US10C\_PUBCOMB.pep:\*
- 16: /cgn2\_6/ptodata/2/pubpaa/US10\_NEW\_PUB.pep:\*
- 17: /cgn2\_6/ptodata/2/pubpaa/US60\_NEW\_PUB.pep:\*
- 18: /cgn2\_6/ptodata/2/pubpaa/US60\_PUBCOMB.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	1745	100.0	317	US-09-946-374-322	Sequence 322, App
2	1745	100.0	317	US-10-147-493-524	Sequence 524, App
3	1745	100.0	317	US-10-145-127-524	Sequence 524, App
4	1745	100.0	317	US-10-160-503-524	Sequence 524, App
5	1745	100.0	317	US-10-143-118-524	Sequence 524, App
6	1745	100.0	317	US-10-144-993-524	Sequence 524, App
7	1745	100.0	317	US-10-158-787-524	Sequence 524, App
8	1745	100.0	317	US-10-008-867-116	Sequence 116, App
9	1745	100.0	317	US-10-063-547-116	Sequence 116, App
10	1745	100.0	317	US-10-028-072-524	Sequence 524, App
11	1745	100.0	317	US-10-063-616-116	Sequence 116, App
12	1745	100.0	317	US-10-121-049-524	Sequence 524, App
13	1745	100.0	317	US-10-123-904-524	Sequence 524, App
14	1745	100.0	317	US-10-140-470-524	Sequence 524, App
15	1745	100.0	317	US-10-063-502-116	Sequence 116, App

16	1745	100.0	317	US-10-175-746-524	Sequence 524, App
17	1745	100.0	317	US-10-176-918-524	Sequence 524, App
18	1745	100.0	317	US-10-176-921-524	Sequence 524, App
19	1745	100.0	317	US-10-137-865-524	Sequence 524, App
20	1745	100.0	317	US-10-140-474-524	Sequence 524, App
21	1745	100.0	317	US-10-142-431-524	Sequence 524, App
22	1745	100.0	317	US-10-143-114-524	Sequence 524, App
23	1745	100.0	317	US-10-140-002-524	Sequence 524, App
24	1745	100.0	317	US-10-006-856A-322	Sequence 322, App
25	1745	100.0	317	US-10-142-419-524	Sequence 524, App
26	1745	100.0	317	US-10-063-518-116	Sequence 116, App
27	1745	100.0	317	US-10-123-262-524	Sequence 524, App
28	1745	100.0	317	US-10-142-423-524	Sequence 524, App
29	1745	100.0	317	US-10-063-598-116	Sequence 116, App
30	1745	100.0	317	US-10-227-693-116	Sequence 322, App
31	1745	100.0	317	US-10-006-818A-322	Sequence 322, App
32	1745	100.0	317	US-10-121-050-524	Sequence 524, App
33	1745	100.0	317	US-10-141-755-524	Sequence 524, App
34	1745	100.0	317	US-10-143-032-524	Sequence 524, App
35	1745	100.0	317	US-10-123-108-524	Sequence 524, App
36	1745	100.0	317	US-10-123-236-524	Sequence 524, App
37	1745	100.0	317	US-10-123-261-524	Sequence 524, App
38	1745	100.0	317	US-10-140-921-524	Sequence 524, App
39	1745	100.0	317	US-10-140-928-524	Sequence 524, App
40	1745	100.0	317	US-10-015-593A-322	Sequence 322, App
41	1745	100.0	317	US-10-063-567-116	Sequence 116, App
42	1745	100.0	317	US-10-015-869A-322	Sequence 322, App
43	1745	100.0	317	US-10-063-538-116	Sequence 116, App
44	1745	100.0	317	US-10-121-045-524	Sequence 524, App
45	1745	100.0	317	US-10-123-292-524	Sequence 524, App

#### ALIGNMENTS

RESULT 1  
US-09-946-374-322  
Sequence 322, Application US/09946374  
Publication No. US20030073129A1  
GENERAL INFORMATION:  
APPLICANT: Baker, Kevin P.  
APPLICANT: Botstein, David  
APPLICANT: Desnoyers, Luc  
APPLICANT: Eaton, Dan L.  
APPLICANT: Ferrara, Napoleone  
APPLICANT: Fong, Sherman  
APPLICANT: Gao, Wei-Qiang  
APPLICANT: Goddard, Audrey  
APPLICANT: Godowski, Paul J.  
APPLICANT: Grimaldi, Christopher J.  
APPLICANT: Gurney, Austin L.  
APPLICANT: Hillan, Kenneth J.  
APPLICANT: Pan, James  
APPLICANT: Paoni, Nicholas F.  
APPLICANT: Roy, Margaret Ann  
APPLICANT: Smith, Victoria  
APPLICANT: Stewart, Timothy A.  
APPLICANT: Tumas, Daniel  
APPLICANT: Watanabe, Colin K.  
APPLICANT: Williams, P. Mickey  
TITLE OF INVENTION: Secreted and Transmembrane Polypeptides and Nucleic  
FILE REFERENCE: P2830P1  
CURRENT APPLICATION NUMBER: US/09/946,374  
PRIOR FILING DATE: 2001-09-04  
PRIOR APPLICATION NUMBER: 60/098716  
PRIOR FILING DATE: 1998-09-01  
PRIOR APPLICATION NUMBER: 60/098723  
PRIOR FILING DATE: 1998-09-01  
PRIOR APPLICATION NUMBER: 60/098749  
PRIOR FILING DATE: 1998-09-01  
PRIOR APPLICATION NUMBER: 60/098750



PRIOR FILING DATE: 1998-10-14  
PRIOR APPLICATION NUMBER: 60/104987  
PRIOR FILING DATE: 1998-10-20  
PRIOR APPLICATION NUMBER: 60/105000  
PRIOR FILING DATE: 1998-10-20  
PRIOR APPLICATION NUMBER: 60/105002  
PRIOR FILING DATE: 1998-10-20  
PRIOR APPLICATION NUMBER: 60/105104  
PRIOR FILING DATE: 1998-10-21  
PRIOR APPLICATION NUMBER: 60/105169  
PRIOR FILING DATE: 1998-10-22  
PRIOR APPLICATION NUMBER: 60/105266  
PRIOR FILING DATE: 1998-10-22  
PRIOR APPLICATION NUMBER: 60/105693  
PRIOR FILING DATE: 1998-10-26  
PRIOR APPLICATION NUMBER: 60/105694  
PRIOR FILING DATE: 1998-10-26  
PRIOR APPLICATION NUMBER: 60/105807

Query Match 100.0%; Score 1745; DB 10; Length 317;  
Best Local Similarity 100.0%; Pred. No. 2.5e-156;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCECHILNAEAFKSKIKCSLKICGVFGIALLTLIVLFMGSKHFWPEVPRK 60  
DB 1 MAKNPENCECHILNAEAFKSKIKCSLKICGVFGIALLTLIVLFMGSKHFWPEVPRK 60  
QY 61 AYMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLVHDPKNGYTGIVFGLOKC 120  
DB 61 AYMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLVHDPKNGYTGIVFGLOKC 120  
QY 121 FIKTOIKVIPSFSEBEIDENEITTTFFEQSVIWPAAKPIENRDFLKNKILEICDN 180  
DB 121 FIKTOIKVIPSFSEBEIDENEITTTFFEQSVIWPAAKPIENRDFLKNKILEICDN 180  
QY 181 VTYWMINPTLISVSELQDFEEBEGDLHFPANEKKGIEONEQWVPQVVEKTRHARQASE 240  
DB 181 VTYWMINPTLISVSELQDFEEBEGDLHFPANEKKGIEONEQWVPQVVEKTRHARQASE 240  
QY 241 BELPINDYTGNGIEBPDMLDERGYCCICRRGNRYCRVCEPILGYPPYCYOGGRVTC 300  
DB 241 BELPINDYTGNGIEBPDMLDERGYCCICRRGNRYCRVCEPILGYPPYCYOGGRVTC 300  
QY 301 RVIMPCNMWVAPMLGRV 317  
DB 301 RVIMPCNMWVAPMLGRV 317

RESULT 2  
US-10-147-493-524  
Sequence 524, Application US/10147493  
Publication No. US2004002921A1  
GENERAL INFORMATION:  
APPLICANT: Baker, Kevin P.  
APPLICANT: Beresini, Maureen  
APPLICANT: Deforge, Laura  
APPLICANT: Desnoyers, Luc  
APPLICANT: Filvaroff, Ellen  
APPLICANT: Gao, Wei-Qiang  
APPLICANT: Gerritsen, Mary E.  
APPLICANT: Goddard, Audrey  
APPLICANT: Godowski, Paul J.  
APPLICANT: Gurney, Austin L.  
APPLICANT: Sherwood, Steven  
APPLICANT: Smith, Victoria  
APPLICANT: Stewart, Timothy A.  
APPLICANT: Tumas, Daniel  
APPLICANT: Watanabe, Colin K  
APPLICANT: Wood, William  
APPLICANT: Zhang, Zemin  
TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC  
FILE REFERENCE: P3330R1C345

CURRENT APPLICATION NUMBER: US/10/147,493  
CURRENT FILING DATE: 2002-05-17  
Prior Application removed - See File Wrapper or Palm  
NUMBER OF SEQ ID NOS: 550  
SEQ ID NO: 524  
LENGTH: 317  
TYPE: PRT  
ORGANISM: Homo Sapien  
US-10-147-493-524

Query Match 100.0%; Score 1745; DB 12; Length 317;  
Best Local Similarity 100.0%; Pred. No. 2.5e-156;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCECHILNAEAFKSKIKCSLKICGVFGIALLTLIVLFMGSKHFWPEVPRK 60  
DB 1 MAKNPENCECHILNAEAFKSKIKCSLKICGVFGIALLTLIVLFMGSKHFWPEVPRK 60  
QY 61 AYMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLVHDPKNGYTGIVFGLOKC 120  
DB 61 AYMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLVHDPKNGYTGIVFGLOKC 120  
QY 121 FIKTOIKVIPSFSEBEIDENEITTTFFEQSVIWPAAKPIENRDFLKNKILEICDN 180  
DB 121 FIKTOIKVIPSFSEBEIDENEITTTFFEQSVIWPAAKPIENRDFLKNKILEICDN 180  
QY 181 VTYWMINPTLISVSELQDFEEBEGDLHFPANEKKGIEONEQWVPQVVEKTRHARQASE 240  
DB 181 VTYWMINPTLISVSELQDFEEBEGDLHFPANEKKGIEONEQWVPQVVEKTRHARQASE 240  
QY 241 BELPINDYTGNGIEBPDMLDERGYCCICRRGNRYCRVCEPILGYPPYCYOGGRVTC 300  
DB 241 BELPINDYTGNGIEBPDMLDERGYCCICRRGNRYCRVCEPILGYPPYCYOGGRVTC 300  
QY 301 RVIMPCNMWVAPMLGRV 317  
DB 301 RVIMPCNMWVAPMLGRV 317

RESULT 3  
US-10-145-127-524  
Sequence 524, Application US/10145127  
Publication No. US2004003558A1  
GENERAL INFORMATION:  
APPLICANT: Baker, Kevin P.  
APPLICANT: Beresini, Maureen  
APPLICANT: Deforge, Laura  
APPLICANT: Desnoyers, Luc  
APPLICANT: Filvaroff, Ellen  
APPLICANT: Gao, Wei-Qiang  
APPLICANT: Gerritsen, Mary E.  
APPLICANT: Goddard, Audrey  
APPLICANT: Godowski, Paul J.  
APPLICANT: Gurney, Austin L.  
APPLICANT: Sherwood, Steven  
APPLICANT: Smith, Victoria  
APPLICANT: Stewart, Timothy A.  
APPLICANT: Tumas, Daniel  
APPLICANT: Watanabe, Colin K  
APPLICANT: Wood, William  
APPLICANT: Zhang, Zemin  
TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC  
FILE REFERENCE: P3330R1C252  
CURRENT APPLICATION NUMBER: US/10/145,127  
CURRENT FILING DATE: 2002-05-13  
Prior Application removed - See File Wrapper or Palm  
NUMBER OF SEQ ID NOS: 550  
SEQ ID NO: 524  
LENGTH: 317  
TYPE: PRT  
ORGANISM: Homo Sapien  
US-10-145-127-524

```

Query Match          100.0%; Score 1745; DB 12; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENEDCHILNAEAFKSKKICSLKICGLVFGILATLLVLPFGSKGFWEVVPK 60
DB 1 MAKNPENEDCHILNAEAFKSKKICSLKICGLVFGILATLLVLPFGSKGFWEVVPK 60
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQKC 120
DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQKC 120
QY 121 FIKTQIKVLPFSEPEEBEIDENEETITTFEESQVIVWPAEKPIENRDLKNSKILIEICDN 180
DB 121 FIKTQIKVLPFSEPEEBEIDENEETITTFEESQVIVWPAEKPIENRDLKNSKILIEICDN 180
QY 181 VTMWINPTLISVSELQDFEEGEDLHFPANEKKGIEQNEQWVWPQVYKTRHARQASE 240
DB 181 VTMWINPTLISVSELQDFEEGEDLHFPANEKKGIEQNEQWVWPQVYKTRHARQASE 240
QY 241 EELPINDYTNGIEFPDMLDERGYCCICRGNRVCRAVCEPILGYPPYCYQSGRVIC 300
DB 241 EELPINDYTNGIEFPDMLDERGYCCICRGNRVCRAVCEPILGYPPYCYQSGRVIC 300
QY 301 RVIMPCNMWVARMGRV 317
DB 301 RVIMPCNMWVARMGRV 317

RESULT 4
US-10-160-503-524
; Sequence 524, Application US/10160503
; Publication No. US20040033559A1
; GENERAL INFORMATION:
; APPLICANT: Baker, Kevin P.
; APPLICANT: Beresini, Maureen
; APPLICANT: Deforge, Laura
; APPLICANT: Desnoyers, Luc
; APPLICANT: Filvaroff, Ellen
; APPLICANT: Gao, Wei-Qiang
; APPLICANT: Gerritsen, Mary E.
; APPLICANT: Goddard, Audrey
; APPLICANT: Godowski, Paul J.
; APPLICANT: Gurney, Austin L.
; APPLICANT: Sherwood, Steven
; APPLICANT: Smith, Victoria
; APPLICANT: Stewart, Timothy A.
; APPLICANT: Tumas, Daniel
; APPLICANT: Watanabe, Colin K
; APPLICANT: Wood, William
; APPLICANT: Zhang, Zemin
; TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
; FILE REFERENCE: P3330R1C446
; CURRENT APPLICATION NUMBER: US/10/160.503
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 550
; SEQ ID NO 524
; LENGTH: 317
; TYPE: PRT
; ORGANISM: Homo Sapien
US-10-160-503-524

Query Match          100.0%; Score 1745; DB 12; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENEDCHILNAEAFKSKKICSLKICGLVFGILATLLVLPFGSKGFWEVVPK 60
DB 1 MAKNPENEDCHILNAEAFKSKKICSLKICGLVFGILATLLVLPFGSKGFWEVVPK 60
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQKC 120

```

```

DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQKC 120
QY 121 FIKTQIKVLPFSEPEEBEIDENEETITTFEESQVIVWPAEKPIENRDLKNSKILIEICDN 180
DB 121 FIKTQIKVLPFSEPEEBEIDENEETITTFEESQVIVWPAEKPIENRDLKNSKILIEICDN 180
QY 181 VTMWINPTLISVSELQDFEEGEDLHFPANEKKGIEQNEQWVWPQVYKTRHARQASE 240
DB 181 VTMWINPTLISVSELQDFEEGEDLHFPANEKKGIEQNEQWVWPQVYKTRHARQASE 240
QY 241 EELPINDYTNGIEFPDMLDERGYCCICRGNRVCRAVCEPILGYPPYCYQSGRVIC 300
DB 241 EELPINDYTNGIEFPDMLDERGYCCICRGNRVCRAVCEPILGYPPYCYQSGRVIC 300
QY 301 RVIMPCNMWVARMGRV 317
DB 301 RVIMPCNMWVARMGRV 317

RESULT 5
US-10-143-118-524
; Sequence 524, Application US/10143118
; Publication No. US20040038335A1
; GENERAL INFORMATION:
; APPLICANT: Baker, Kevin P.
; APPLICANT: Beresini, Maureen
; APPLICANT: Deforge, Laura
; APPLICANT: Desnoyers, Luc
; APPLICANT: Filvaroff, Ellen
; APPLICANT: Gao, Wei-Qiang
; APPLICANT: Gerritsen, Mary E.
; APPLICANT: Goddard, Audrey
; APPLICANT: Godowski, Paul J.
; APPLICANT: Gurney, Austin L.
; APPLICANT: Sherwood, Steven
; APPLICANT: Smith, Victoria
; APPLICANT: Stewart, Timothy A.
; APPLICANT: Tumas, Daniel
; APPLICANT: Watanabe, Colin K
; APPLICANT: Wood, William
; APPLICANT: Zhang, Zemin
; TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
; FILE REFERENCE: P3330R1C228
; CURRENT APPLICATION NUMBER: US/10/143.118
; CURRENT FILING DATE: 2002-05-09
; Prior Application removed - See Palm or File Wrapper
; NUMBER OF SEQ ID NOS: 550
; SEQ ID NO 524
; LENGTH: 317
; TYPE: PRT
; ORGANISM: Homo Sapien
US-10-143-118-524

Query Match          100.0%; Score 1745; DB 12; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENEDCHILNAEAFKSKKICSLKICGLVFGILATLLVLPFGSKGFWEVVPK 60
DB 1 MAKNPENEDCHILNAEAFKSKKICSLKICGLVFGILATLLVLPFGSKGFWEVVPK 60
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQKC 120
DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIFVGLQKC 120
QY 121 FIKTQIKVLPFSEPEEBEIDENEETITTFEESQVIVWPAEKPIENRDLKNSKILIEICDN 180
DB 121 FIKTQIKVLPFSEPEEBEIDENEETITTFEESQVIVWPAEKPIENRDLKNSKILIEICDN 180
QY 181 VTMWINPTLISVSELQDFEEGEDLHFPANEKKGIEQNEQWVWPQVYKTRHARQASE 240

```

```
Db      181 VTMWNPILISVSEIQDFEEBEGEDLHPANEKKGIQNEQWVVPQVYKTRHARQASE 240
Qy      241 EELPINDYENGIEPDMLEDERGYCCICRGNRRCRVCEPILGYPPYCYOGGRVLC 300
Db      241 EELPINDYENGIEPDMLEDERGYCCICRGNRRCRVCEPILGYPPYCYOGGRVLC 300
Qy      301 RVIMPCNMWVARMUGRV 317
Db      301 RVIMPCNMWVARMUGRV 317

RESULT 6
US-10-144-993-524
; Sequence 524, Application US/10144993
; Publication No. US2004003836A1
; GENERAL INFORMATION:
; APPLICANT: Baker, Kevin P.
; APPLICANT: Beresini, Maureen
; APPLICANT: DeForge, Laura
; APPLICANT: Desnoyers, Luc
; APPLICANT: Filvaroff, Ellen
; APPLICANT: Gao, Wei-Qiang
; APPLICANT: Gerritsen, Mary E.
; APPLICANT: Goddard, Audrey
; APPLICANT: Godowski, Paul J.
; APPLICANT: Gurney, Austin L.
; APPLICANT: Sherwood, Steven
; APPLICANT: Smith, Victoria
; APPLICANT: Stewart, Timothy A.
; APPLICANT: Tumas, Daniel
; APPLICANT: Watanabe, Colin K
; APPLICANT: Wood, William
; APPLICANT: Zhang, Zemin
; TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
; FILE OF INVENTION: ACIDS ENCODING THE SAME
; FILE REFERENCE: P330R1C261
; CURRENT APPLICATION NUMBER: US/10/144, 993
; CURRENT FILING DATE: 2002-05-13
; Prior Application removed - See File Wrapper or Palm
; NUMBER OF SEQ ID NOS: 550
; SEQ ID NO 524
; LENGTH: 317
; TYPE: PRT
; ORGANISM: Homo Sapien
US-10-144-993-524

Query Match      100.0%; Score 1745; DB 12; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156; Indels 0; Gaps 0;
Matches 317; Conservative 0; Mismatches 0;

Qy      1  MAKNPENCEDCHILNAEAFKSKKICKSLKICGLVFGILATLILVFWGSKHFWEVPKX 60
Db      1  MAKNPENCEDCHILNAEAFKSKKICKSLKICGLVFGILATLILVFWGSKHFWEVPKX 60
Qy      61 AYMEHTFYSNGEKKKIYMEIDPVTREIFRSNGTDETLVHDFKNGYTGIVFGLOXC 120
Db      61 AYMEHTFYSNGEKKKIYMEIDPVTREIFRSNGTDETLVHDFKNGYTGIVFGLOXC 120
Qy      121 FIKTOIKVIPFSPPEEIDENEIITTTFFEQSVIWPAPKPIENRDLKSKLIEICDN 180
Db      121 FIKTOIKVIPFSPPEEIDENEIITTTFFEQSVIWPAPKPIENRDLKSKLIEICDN 180
Qy      181 VTMWNPILISVSEIQDFEEBEGEDLHPANEKKGIQNEQWVVPQVYKTRHARQASE 240
Db      181 VTMWNPILISVSEIQDFEEBEGEDLHPANEKKGIQNEQWVVPQVYKTRHARQASE 240
Qy      241 EELPINDYENGIEPDMLEDERGYCCICRGNRRCRVCEPILGYPPYCYOGGRVLC 300
Db      241 EELPINDYENGIEPDMLEDERGYCCICRGNRRCRVCEPILGYPPYCYOGGRVLC 300
Qy      301 RVIMPCNMWVARMUGRV 317
Db      301 RVIMPCNMWVARMUGRV 317
```

```
RESULT 7
US-10-158-787-524
; Sequence 524, Application US/10158787
; Publication No. US20040039164A1
; GENERAL INFORMATION:
; APPLICANT: Baker, Kevin P.
; APPLICANT: Beresini, Maureen
; APPLICANT: DeForge, Laura
; APPLICANT: Desnoyers, Luc
; APPLICANT: Filvaroff, Ellen
; APPLICANT: Gao, Wei-Qiang
; APPLICANT: Gerritsen, Mary E.
; APPLICANT: Goddard, Audrey
; APPLICANT: Godowski, Paul J.
; APPLICANT: Gurney, Austin L.
; APPLICANT: Sherwood, Steven
; APPLICANT: Smith, Victoria
; APPLICANT: Stewart, Timothy A.
; APPLICANT: Tumas, Daniel
; APPLICANT: Watanabe, Colin K
; APPLICANT: Wood, William
; APPLICANT: Zhang, Zemin
; TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
; FILE OF INVENTION: ACIDS ENCODING THE SAME
; FILE REFERENCE: P330R1C49
; CURRENT APPLICATION NUMBER: US/10/158, 787
; CURRENT FILING DATE: 2003-04-03
; PRIOR APPLICATION NUMBER: 60/049911
; PRIOR FILING DATE: 1997-06-18
; PRIOR APPLICATION NUMBER: 60/056974
; PRIOR FILING DATE: 1997-08-26
; PRIOR APPLICATION NUMBER: 60/059113
; PRIOR FILING DATE: 1997-09-17
; PRIOR APPLICATION NUMBER: 60/059115
; PRIOR FILING DATE: 1997-09-17
; PRIOR APPLICATION NUMBER: 60/059117
; PRIOR FILING DATE: 1997-09-17
; PRIOR APPLICATION NUMBER: 60/059122
; PRIOR FILING DATE: 1997-09-17
; PRIOR APPLICATION NUMBER: 60/059184
; PRIOR FILING DATE: 1997-09-17
; PRIOR APPLICATION NUMBER: 60/059263
; PRIOR FILING DATE: 1997-09-18
; PRIOR APPLICATION NUMBER: 60/059352
; PRIOR FILING DATE: 1997-09-19
; PRIOR APPLICATION NUMBER: 60/059588
; PRIOR FILING DATE: 1997-09-19
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 550
; SEQ ID NO 524
; LENGTH: 317
; TYPE: PRT
; ORGANISM: Homo Sapien
US-10-158-787-524

Query Match      100.0%; Score 1745; DB 12; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156; Indels 0; Gaps 0;
Matches 317; Conservative 0; Mismatches 0;

Qy      1  MAKNPENCEDCHILNAEAFKSKKICKSLKICGLVFGILATLILVFWGSKHFWEVPKX 60
Db      1  MAKNPENCEDCHILNAEAFKSKKICKSLKICGLVFGILATLILVFWGSKHFWEVPKX 60
Qy      61 AYMEHTFYSNGEKKKIYMEIDPVTREIFRSNGTDETLVHDFKNGYTGIVFGLOXC 120
Db      61 AYMEHTFYSNGEKKKIYMEIDPVTREIFRSNGTDETLVHDFKNGYTGIVFGLOXC 120
Qy      121 FIKTOIKVIPFSPPEEIDENEIITTTFFEQSVIWPAPKPIENRDLKSKLIEICDN 180
Db      121 FIKTOIKVIPFSPPEEIDENEIITTTFFEQSVIWPAPKPIENRDLKSKLIEICDN 180
```

QY 181 UTMWNTPLISVSELODEEEDGDLHPANERKGIENBQWVQVYVETKRRAROSE 240  
DB 181 UTMWNTPLISVSELODEEEDGDLHPANERKGIENBQWVQVYVETKRRAROSE 240  
QY 241 EELPINDYENGIEFDPMDERGYCCICRGRNRCRYCEPLTGYYPYCYOGGRVTC 300  
DB 241 EELPINDYENGIEFDPMDERGYCCICRGRNRCRYCEPLTGYYPYCYOGGRVTC 300  
QY 301 RYIMPCNMVWVAMTGRV 317  
DB 301 RYIMPCNMVWVAMTGRV 317  
RESULT 8  
US-10-006-867-116  
Sequence 116, Application US/10006867  
Publication No. US20020119130A1  
GENERAL INFORMATION:  
APPLICANT: Eaton, Dan L.  
APPLICANT: Filvaroff, Ellen  
APPLICANT: Geritsen, Mary E.  
APPLICANT: Goddard, Audrey  
APPLICANT: Godowski, Paul J.  
APPLICANT: Grimaldi, Christopher J.  
APPLICANT: Guiney, Austin L.  
APPLICANT: Watanabe, Colin K.  
APPLICANT: Wood, William I.  
TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC  
FILE OF INVENTION: ACIDS ENCODING THE SAME  
FILE REFERENCE: P3320R1C1  
CURRENT APPLICATION NUMBER: US/10/006,867  
CURRENT FILING DATE: 2001-12-06  
PRIOR APPLICATION NUMBER: 60/063435  
PRIOR FILING DATE: 1997-10-29  
PRIOR APPLICATION NUMBER: 60/064215  
PRIOR FILING DATE: 1997-10-29  
PRIOR APPLICATION NUMBER: 60/082797  
PRIOR FILING DATE: 1998-04-22  
PRIOR APPLICATION NUMBER: 60/083495  
PRIOR FILING DATE: 1998-04-29  
PRIOR APPLICATION NUMBER: 60/085579  
PRIOR FILING DATE: 1998-05-15  
PRIOR APPLICATION NUMBER: 60/087759  
PRIOR FILING DATE: 1998-06-02  
PRIOR APPLICATION NUMBER: 60/088021  
PRIOR FILING DATE: 1998-06-04  
PRIOR APPLICATION NUMBER: 60/088029  
PRIOR FILING DATE: 1998-06-04  
PRIOR APPLICATION NUMBER: 60/088030  
PRIOR FILING DATE: 1998-06-04  
PRIOR APPLICATION NUMBER: 60/088734  
PRIOR FILING DATE: 1998-06-10  
PRIOR APPLICATION NUMBER: 60/088740  
PRIOR FILING DATE: 1998-06-10  
PRIOR APPLICATION NUMBER: 60/088811  
PRIOR FILING DATE: 1998-06-10  
PRIOR APPLICATION NUMBER: 60/088824  
PRIOR FILING DATE: 1998-06-10  
PRIOR APPLICATION NUMBER: 60/088825  
PRIOR FILING DATE: 1998-06-10  
PRIOR APPLICATION NUMBER: 60/088863  
PRIOR FILING DATE: 1998-06-11  
PRIOR APPLICATION NUMBER: 60/089105  
PRIOR FILING DATE: 1998-06-12  
PRIOR APPLICATION NUMBER: 60/089514  
PRIOR FILING DATE: 1998-06-16  
PRIOR APPLICATION NUMBER: 60/089653  
PRIOR FILING DATE: 1998-06-17  
PRIOR APPLICATION NUMBER: 60/089952  
PRIOR FILING DATE: 1998-06-19  
PRIOR APPLICATION NUMBER: 60/090246  
PRIOR FILING DATE: 1998-06-22  
PRIOR APPLICATION NUMBER: 60/090444  
PRIOR FILING DATE: 1998-06-24  
PRIOR APPLICATION NUMBER: 60/090688  
PRIOR FILING DATE: 1998-06-25  
PRIOR APPLICATION NUMBER: 60/090696  
PRIOR FILING DATE: 1998-06-25  
PRIOR APPLICATION NUMBER: 60/090862  
PRIOR FILING DATE: 1998-06-26  
PRIOR APPLICATION NUMBER: 60/091628  
PRIOR FILING DATE: 1998-07-02  
PRIOR APPLICATION NUMBER: 60/096012  
PRIOR FILING DATE: 1998-08-10  
PRIOR APPLICATION NUMBER: 60/096757  
PRIOR FILING DATE: 1998-08-17  
PRIOR APPLICATION NUMBER: 60/096949  
PRIOR FILING DATE: 1998-08-18  
PRIOR APPLICATION NUMBER: 60/096959  
PRIOR FILING DATE: 1998-08-18  
PRIOR APPLICATION NUMBER: 60/097954  
PRIOR FILING DATE: 1998-08-26  
PRIOR APPLICATION NUMBER: 60/097971  
PRIOR FILING DATE: 1998-08-26  
PRIOR APPLICATION NUMBER: 60/097979  
PRIOR FILING DATE: 1998-08-26  
PRIOR APPLICATION NUMBER: 60/098749  
PRIOR FILING DATE: 1998-09-01  
PRIOR APPLICATION NUMBER: 60/099741  
PRIOR FILING DATE: 1998-09-10  
PRIOR APPLICATION NUMBER: 60/099763  
PRIOR FILING DATE: 1998-09-10  
PRIOR APPLICATION NUMBER: 60/099792  
PRIOR FILING DATE: 1998-09-10  
PRIOR APPLICATION NUMBER: 60/099812  
PRIOR FILING DATE: 1998-09-10  
PRIOR APPLICATION NUMBER: 60/099815  
PRIOR FILING DATE: 1998-09-10  
PRIOR APPLICATION NUMBER: 60/100627  
PRIOR FILING DATE: 1998-09-16  
PRIOR APPLICATION NUMBER: 60/100662  
PRIOR FILING DATE: 1998-09-16  
PRIOR APPLICATION NUMBER: 60/100683  
PRIOR FILING DATE: 1998-09-17  
PRIOR APPLICATION NUMBER: 60/100684  
PRIOR FILING DATE: 1998-09-17  
PRIOR APPLICATION NUMBER: 60/100930  
PRIOR FILING DATE: 1998-09-17  
PRIOR APPLICATION NUMBER: 60/101279  
PRIOR FILING DATE: 1998-09-22  
PRIOR APPLICATION NUMBER: 60/101475  
PRIOR FILING DATE: 1998-09-23  
PRIOR APPLICATION NUMBER: 60/101738  
PRIOR FILING DATE: 1998-09-24  
PRIOR APPLICATION NUMBER: 60/101743  
PRIOR FILING DATE: 1998-09-24  
PRIOR APPLICATION NUMBER: 60/101916  
PRIOR FILING DATE: 1998-09-24  
PRIOR APPLICATION NUMBER: 60/102570  
PRIOR FILING DATE: 1998-09-30  
PRIOR APPLICATION NUMBER: 60/103449  
PRIOR FILING DATE: 1998-10-06  
PRIOR APPLICATION NUMBER: 60/103678  
PRIOR FILING DATE: 1998-10-08  
PRIOR APPLICATION NUMBER: 60/103679  
PRIOR FILING DATE: 1998-10-08  
PRIOR APPLICATION NUMBER: 60/103711  
PRIOR FILING DATE: 1998-10-08  
PRIOR APPLICATION NUMBER: 60/105000  
PRIOR FILING DATE: 1998-10-20  
PRIOR APPLICATION NUMBER: 60/105002  
PRIOR FILING DATE: 1998-10-20  
PRIOR APPLICATION NUMBER: 60/105881  
PRIOR FILING DATE: 1998-10-27  
PRIOR APPLICATION NUMBER: 60/106030  
PRIOR FILING DATE: 1998-10-28

PRIOR APPLICATION NUMBER: 60/106464  
 PRIOR FILING DATE: 1998-10-30  
 PRIOR APPLICATION NUMBER: 60/106856  
 PRIOR FILING DATE: 1998-11-03  
 PRIOR APPLICATION NUMBER: 60/108807  
 PRIOR FILING DATE: 1998-11-17  
 PRIOR APPLICATION NUMBER: 60/112419  
 PRIOR FILING DATE: 1998-12-15  
 PRIOR APPLICATION NUMBER: 60/112422  
 PRIOR FILING DATE: 1998-12-15  
 PRIOR APPLICATION NUMBER: 60/112853  
 PRIOR FILING DATE: 1998-12-16  
 PRIOR APPLICATION NUMBER: 60/113011  
 PRIOR FILING DATE: 1998-12-16  
 PRIOR APPLICATION NUMBER: 60/112854  
 PRIOR FILING DATE: 1998-12-16  
 PRIOR APPLICATION NUMBER: 60/113300  
 PRIOR FILING DATE: 1998-12-22  
 PRIOR APPLICATION NUMBER: 60/113408  
 PRIOR FILING DATE: 1998-12-22  
 PRIOR APPLICATION NUMBER: 60/113430  
 PRIOR FILING DATE: 1998-12-23  
 PRIOR APPLICATION NUMBER: 60/113621  
 PRIOR FILING DATE: 1998-12-23  
 PRIOR APPLICATION NUMBER: 60/114223  
 PRIOR FILING DATE: 1998-12-30  
 PRIOR APPLICATION NUMBER: 60/115614  
 PRIOR FILING DATE: 1999-01-12  
 PRIOR APPLICATION NUMBER: 60/116527  
 PRIOR FILING DATE: 1999-01-20  
 PRIOR APPLICATION NUMBER: 60/116843  
 PRIOR FILING DATE: 1999-01-22  
 PRIOR APPLICATION NUMBER: 60/119285  
 PRIOR FILING DATE: 1999-02-09  
 PRIOR APPLICATION NUMBER: 60/119287  
 PRIOR FILING DATE: 1999-02-09  
 PRIOR APPLICATION NUMBER: 60/119525  
 PRIOR FILING DATE: 1999-02-10  
 PRIOR APPLICATION NUMBER: 60/119549  
 PRIOR FILING DATE: 1999-02-10  
 PRIOR APPLICATION NUMBER: 60/120014  
 PRIOR FILING DATE: 1999-02-11  
 PRIOR APPLICATION NUMBER: 60/129122  
 PRIOR FILING DATE: 1999-04-13  
 PRIOR APPLICATION NUMBER: 60/129674  
 PRIOR FILING DATE: 1999-04-16  
 PRIOR APPLICATION NUMBER: 60/131291  
 PRIOR FILING DATE: 1999-04-27  
 PRIOR APPLICATION NUMBER: 60/138387  
 PRIOR FILING DATE: 1999-06-09  
 PRIOR APPLICATION NUMBER: 60/144791  
 PRIOR FILING DATE: 1999-07-20  
 PRIOR APPLICATION NUMBER: 60/169495  
 PRIOR FILING DATE: 1999-12-07  
 PRIOR APPLICATION NUMBER: 60/175481  
 PRIOR FILING DATE: 2000-01-11  
 PRIOR APPLICATION NUMBER: 60/191007  
 PRIOR FILING DATE: 2000-03-21  
 PRIOR APPLICATION NUMBER: 60/199397  
 PRIOR FILING DATE: 2000-04-25  
 PRIOR APPLICATION NUMBER: 09/380139  
 PRIOR FILING DATE: 1998-08-25  
 PRIOR APPLICATION NUMBER: 09/311832  
 PRIOR FILING DATE: 1999-05-14  
 PRIOR APPLICATION NUMBER: 09/380137  
 PRIOR FILING DATE: 1999-08-25  
 PRIOR APPLICATION NUMBER: 09/380138  
 PRIOR FILING DATE: 1999-08-25  
 PRIOR APPLICATION NUMBER: 09/380142

Query Match 100.0%; Score 1745; DB 13; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 2.5e-156;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCEDCHITINAAFKSKIKCSLKTGCVFGIATLTLVLFMGSKHFMPEVPEKK 60  
 DB 1 MAKNPENCEDCHITINAAFKSKIKCSLKTGCVFGIATLTLVLFMGSKHFMPEVPEKK 60  
 QY 61 AYDMHTFYSNGEKKKIYMEIDPVTREIFPSGNGTDETLVHDFKNGYTGIFVGLQKC 120  
 DB 61 AYDMHTFYSNGEKKKIYMEIDPVTREIFPSGNGTDETLVHDFKNGYTGIFVGLQKC 120  
 QY 121 FIKTOIKVIPSESEEBEIDNEETTTFFQSVIYWAEPKPIENRDLKNSKILIEDN 180  
 DB 121 FIKTOIKVIPSESEEBEIDNEETTTFFQSVIYWAEPKPIENRDLKNSKILIEDN 180  
 QY 121 FIKTOIKVIPSESEEBEIDNEETTTFFQSVIYWAEPKPIENRDLKNSKILIEDN 180  
 DB 121 FIKTOIKVIPSESEEBEIDNEETTTFFQSVIYWAEPKPIENRDLKNSKILIEDN 180  
 QY 181 VTMWINFTLISVSELQDFEEGEDLHPANEEKGIEQNEQWVVPQVVEKTRHARQASE 240  
 DB 181 VTMWINFTLISVSELQDFEEGEDLHPANEEKGIEQNEQWVVPQVVEKTRHARQASE 240  
 QY 241 BELPINDYTENGIEDPMLDERGYCICRGGRNRCRVCEPPLIGYPIYPCYGGRVIC 300  
 DB 241 BELPINDYTENGIEDPMLDERGYCICRGGRNRCRVCEPPLIGYPIYPCYGGRVIC 300  
 QY 301 RYIMPCNMWVARMIGRV 317  
 DB 301 RYIMPCNMWVARMIGRV 317

RESULT 9  
 US-10-063-547-116  
 Sequence 116, Application US/10063547  
 Publication No. US20020182638A1  
 GENERAL INFORMATION:  
 APPLICANT: Baton, Dan L.  
 APPLICANT: Filvaroff, Ellen  
 APPLICANT: Gerriksen, Mary E.  
 APPLICANT: Goddard, Audrey  
 APPLICANT: Godowski, Paul J.  
 APPLICANT: Grimaldi, Christopher J.  
 APPLICANT: Gurney, Austin L.  
 APPLICANT: Matarade, Colin K.  
 APPLICANT: Wood, William I.  
 TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC  
 FILE REFERENCE: P3230R1C1  
 CURRENT APPLICATION NUMBER: US/10/063,547  
 PRIOR FILING DATE: 2002-05-02  
 PRIOR Application removed - See file Wrapper or Palm  
 NUMBER OF SEQ ID NOS: 170  
 SEQ ID NO 116  
 LENGTH: 317  
 TYPE: PRT  
 ORGANISM: Homo Sapien  
 US-10-063-547-116

Query Match 100.0%; Score 1745; DB 13; Length 317;  
 Best Local Similarity 100.0%; Pred. No. 2.5e-156;  
 Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCEDCHITINAAFKSKIKCSLKTGCVFGIATLTLVLFMGSKHFMPEVPEKK 60  
 DB 1 MAKNPENCEDCHITINAAFKSKIKCSLKTGCVFGIATLTLVLFMGSKHFMPEVPEKK 60  
 QY 61 AYDMHTFYSNGEKKKIYMEIDPVTREIFPSGNGTDETLVHDFKNGYTGIFVGLQKC 120  
 DB 61 AYDMHTFYSNGEKKKIYMEIDPVTREIFPSGNGTDETLVHDFKNGYTGIFVGLQKC 120  
 QY 121 FIKTOIKVIPSESEEBEIDNEETTTFFQSVIYWAEPKPIENRDLKNSKILIEDN 180  
 DB 121 FIKTOIKVIPSESEEBEIDNEETTTFFQSVIYWAEPKPIENRDLKNSKILIEDN 180  
 QY 181 VTMWINFTLISVSELQDFEEGEDLHPANEEKGIEQNEQWVVPQVVEKTRHARQASE 240  
 DB 181 VTMWINFTLISVSELQDFEEGEDLHPANEEKGIEQNEQWVVPQVVEKTRHARQASE 240

QY 241 EELPINDYENGIEPDMIDERGYCICYRGRNRYCRVCEBLLGYPPYCYOGGRVYC 300  
DB 241 EELPINDYENGIEPDMIDERGYCICYRGRNRYCRVCEBLLGYPPYCYOGGRVYC 300  
QY 301 RVIMPONMVAARMLGRV 317  
DB 301 RVIMPONMVAARMLGRV 317

RESULT 10  
US-10-028-072-524  
; Sequence 524, Application US/10028072  
; Publication No. US20030004311A1  
; GENERAL INFORMATION:  
; APPLICANT: Baker, Kevin P.  
; APPLICANT: Beresini, Maureen  
; APPLICANT: DeForge, Laura  
; APPLICANT: Desnoyers, Luc  
; APPLICANT: Filvaroff, Ellen  
; APPLICANT: Gao, Wei Qiang  
; APPLICANT: Gerritsen, Mary E.  
; APPLICANT: Goddard, Audrey  
; APPLICANT: Godowski, Paul J.  
; APPLICANT: Gurney, Austin L.  
; APPLICANT: Sherwood, Steven  
; APPLICANT: Smith, Victoria  
; APPLICANT: Stewart, Timothy A.  
; APPLICANT: Tumas, Daniel  
; APPLICANT: Watanabe, Colin K  
; APPLICANT: Wood, William  
; APPLICANT: Zhang  
; TITLE OF INVENTION:  
; FILE REFERENCE:  
; CURRENT APPLICATION NUMBER: US/10/028,072  
; PRIOR FILING DATE: 2001-12-19  
; PRIOR APPLICATION NUMBER: 60/049911  
; PRIOR FILING DATE: 1997-06-18  
; PRIOR APPLICATION NUMBER: 60/056974  
; PRIOR FILING DATE: 1997-08-26  
; PRIOR APPLICATION NUMBER: 60/059113  
; PRIOR FILING DATE: 1997-09-17  
; PRIOR APPLICATION NUMBER: 60/059115  
; PRIOR FILING DATE: 1997-09-17  
; PRIOR APPLICATION NUMBER: 60/059117  
; PRIOR FILING DATE: 1997-09-17  
; PRIOR APPLICATION NUMBER: 60/059122  
; PRIOR FILING DATE: 1997-09-17  
; PRIOR APPLICATION NUMBER: 60/059184  
; PRIOR FILING DATE: 1997-09-17  
; PRIOR APPLICATION NUMBER: 60/059263  
; PRIOR FILING DATE: 1997-09-18  
; PRIOR APPLICATION NUMBER: 60/059352  
; PRIOR FILING DATE: 1997-09-19  
; PRIOR APPLICATION NUMBER: 60/059588  
; PRIOR FILING DATE: 1997-09-19  
; PRIOR APPLICATION NUMBER: 60/059836  
; PRIOR FILING DATE: 1997-09-24  
; PRIOR APPLICATION NUMBER: 60/062250  
; PRIOR FILING DATE: 1997-10-17  
; PRIOR APPLICATION NUMBER: 60/062285  
; PRIOR FILING DATE: 1997-10-17  
; PRIOR APPLICATION NUMBER: 60/062287  
; PRIOR FILING DATE: 1997-10-17  
; PRIOR APPLICATION NUMBER: 60/062814  
; PRIOR FILING DATE: 1997-10-24  
; PRIOR APPLICATION NUMBER: 60/062816  
; PRIOR FILING DATE: 1997-10-24  
; PRIOR APPLICATION NUMBER: 60/063045  
; PRIOR FILING DATE: 1997-10-24  
; PRIOR APPLICATION NUMBER: 60/063082  
; PRIOR FILING DATE: 1997-10-31  
; PRIOR APPLICATION NUMBER: 60/063127  
; PRIOR FILING DATE: 1997-10-24

PRIOR APPLICATION NUMBER: 60/063327  
PRIOR FILING DATE: 1997-10-27  
PRIOR APPLICATION NUMBER: 60/063329  
PRIOR FILING DATE: 1997-10-27  
PRIOR APPLICATION NUMBER: 60/063550  
PRIOR FILING DATE: 1997-10-28  
PRIOR APPLICATION NUMBER: 60/063561  
PRIOR FILING DATE: 1997-10-28  
PRIOR APPLICATION NUMBER: 60/063704  
PRIOR FILING DATE: 1997-10-29  
PRIOR APPLICATION NUMBER: 60/063733  
PRIOR FILING DATE: 1997-10-29  
PRIOR APPLICATION NUMBER: 60/063735  
PRIOR FILING DATE: 1997-10-29  
PRIOR APPLICATION NUMBER: 60/063738  
PRIOR FILING DATE: 1997-10-29  
PRIOR APPLICATION NUMBER: 60/063755  
PRIOR FILING DATE: 1997-10-17  
PRIOR APPLICATION NUMBER: 60/064248  
PRIOR FILING DATE: 1997-11-03  
PRIOR APPLICATION NUMBER: 60/064809  
PRIOR FILING DATE: 1997-11-07  
PRIOR APPLICATION NUMBER: 60/065186  
PRIOR FILING DATE: 1997-11-12  
PRIOR APPLICATION NUMBER: 60/065846  
PRIOR FILING DATE: 1997-11-17  
PRIOR APPLICATION NUMBER: 60/066364  
PRIOR FILING DATE: 1997-11-21  
PRIOR APPLICATION NUMBER: 60/066453  
PRIOR FILING DATE: 1997-11-24  
PRIOR APPLICATION NUMBER: 60/066511  
PRIOR FILING DATE: 1997-11-24  
PRIOR APPLICATION NUMBER: 60/066770  
PRIOR FILING DATE: 1997-11-24  
PRIOR APPLICATION NUMBER: 60/069212  
PRIOR FILING DATE: 1997-12-11  
PRIOR APPLICATION NUMBER: 60/069278  
PRIOR FILING DATE: 1997-12-11  
PRIOR APPLICATION NUMBER: 60/069334  
PRIOR FILING DATE: 1997-12-11  
PRIOR APPLICATION NUMBER: 60/069694  
PRIOR FILING DATE: 1997-12-16  
PRIOR APPLICATION NUMBER: 60/072320  
PRIOR FILING DATE: 1998-01-23  
PRIOR APPLICATION NUMBER: 60/073512  
PRIOR FILING DATE: 1998-02-04  
PRIOR APPLICATION NUMBER: 60/074086  
PRIOR FILING DATE: 1998-02-09  
PRIOR APPLICATION NUMBER: 60/074092  
PRIOR FILING DATE: 1998-02-09  
PRIOR APPLICATION NUMBER: 60/077791  
PRIOR FILING DATE: 1998-03-12  
PRIOR APPLICATION NUMBER: 60/078910  
PRIOR FILING DATE: 1998-03-20  
PRIOR APPLICATION NUMBER: 60/079294  
PRIOR FILING DATE: 1998-03-25  
PRIOR APPLICATION NUMBER: 60/079663  
PRIOR FILING DATE: 1998-02-27  
PRIOR APPLICATION NUMBER: 60/079728  
PRIOR FILING DATE: 1998-03-27  
PRIOR APPLICATION NUMBER: 60/080165  
PRIOR FILING DATE: 1998-03-31  
PRIOR APPLICATION NUMBER: 60/081203  
PRIOR FILING DATE: 1998-04-09  
PRIOR APPLICATION NUMBER: 60/081229  
PRIOR FILING DATE: 1998-04-09  
PRIOR APPLICATION NUMBER: 60/081695  
PRIOR FILING DATE: 1998-04-14  
PRIOR APPLICATION NUMBER: 60/081817  
PRIOR FILING DATE: 1998-04-15  
PRIOR APPLICATION NUMBER: 60/081818  
PRIOR FILING DATE: 1998-04-15  
PRIOR APPLICATION NUMBER: 60/082999



```

PRIOR FILING DATE: 1998-04-24
PRIOR APPLICATION NUMBER: 60/083322
PRIOR FILING DATE: 1998-04-28
PRIOR APPLICATION NUMBER: 60/083545
PRIOR FILING DATE: 1998-04-29
PRIOR APPLICATION NUMBER: 60/084600
PRIOR FILING DATE: 1998-05-07
PRIOR APPLICATION NUMBER: 60/084627
PRIOR FILING DATE: 1998-05-07
PRIOR APPLICATION NUMBER: 60/084637
PRIOR FILING DATE: 1998-05-07
PRIOR APPLICATION NUMBER: 60/085149
PRIOR FILING DATE: 1998-05-12
PRIOR APPLICATION NUMBER: 60/085323
PRIOR FILING DATE: 1998-05-13
PRIOR APPLICATION NUMBER: 60/085338
PRIOR FILING DATE: 1998-05-13
PRIOR APPLICATION NUMBER: 60/085339
PRIOR FILING DATE: 1998-05-13
PRIOR APPLICATION NUMBER: 60/085579
PRIOR FILING DATE: 1998-05-15
PRIOR APPLICATION NUMBER: 60/085697
PRIOR FILING DATE: 1998-05-15
PRIOR APPLICATION NUMBER: 60/085704
PRIOR FILING DATE: 1998-05-15
PRIOR APPLICATION NUMBER: 60/086414
PRIOR FILING DATE: 1998-05-22
PRIOR APPLICATION NUMBER: 60/086430
PRIOR FILING DATE: 1998-05-22
PRIOR APPLICATION NUMBER: 60/087106
PRIOR FILING DATE: 1998-05-28
PRIOR APPLICATION NUMBER: 60/088026
PRIOR FILING DATE: 1998-06-04
PRIOR APPLICATION NUMBER: 60/088730
PRIOR FILING DATE: 1998-06-10
PRIOR APPLICATION NUMBER: 60/088741
PRIOR FILING DATE: 1998-06-10
PRIOR APPLICATION NUMBER: 60/088810
PRIOR FILING DATE: 1998-06-10
PRIOR APPLICATION NUMBER: 60/088858
PRIOR FILING DATE: 1998-06-11
PRIOR APPLICATION NUMBER: 60/089532
PRIOR FILING DATE: 1998-06-17
PRIOR APPLICATION NUMBER: 60/089599
PRIOR FILING DATE: 1998-06-17
PRIOR APPLICATION NUMBER: 60/089907
PRIOR FILING DATE: 1998-06-18
PRIOR APPLICATION NUMBER: 60/089947
PRIOR FILING DATE: 1998-06-19
PRIOR APPLICATION NUMBER: 60/090349
PRIOR FILING DATE: 1998-06-23
PRIOR APPLICATION NUMBER: 60/090429
PRIOR FILING DATE: 1998-06-24
PRIOR APPLICATION NUMBER: 60/090445
PRIOR FILING DATE: 1998-06-24
PRIOR APPLICATION NUMBER: 60/090538
PRIOR FILING DATE: 1998-06-24
PRIOR APPLICATION NUMBER: 60/090863
PRIOR FILING DATE: 1998-06-26
PRIOR APPLICATION NUMBER: 60/091360
PRIOR FILING DATE: 1998-07-01
PRIOR APPLICATION NUMBER: 60/091519
PRIOR FILING DATE: 1998-07-02
PRIOR APPLICATION NUMBER: 60/091982
PRIOR FILING DATE: 1998-07-07

```

```

Query Match 100.0%; Score 1745; DB 14; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
1 MAKNPENCDCIILNAEAFKSKIKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
1 MAKNPENCDCIILNAEAFKSKIKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
1 MAKNPENCDCIILNAEAFKSKIKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60

```

```

QY 61 AYDMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLBVDHFKNGYTGIVFGLQKC 120
DB 61 AYDMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLBVDHFKNGYTGIVFGLQKC 120
QY 121 FIKTOIVIPFSPSEEBEIDENEITTTFFEOQYIWPAPKPIENRFLKNSKILEICDN 180
DB 121 FIKTOIVIPFSPSEEBEIDENEITTTFFEOQYIWPAPKPIENRFLKNSKILEICDN 180
QY 181 VTWYINPTLISVSELODFEPEGEDLHFPANEKKGIEQNEQWVVPQYKVEKTRHARQASE 240
DB 181 VTWYINPTLISVSELODFEPEGEDLHFPANEKKGIEQNEQWVVPQYKVEKTRHARQASE 240
QY 241 EELPINDYNTENGIEFDPMLDERGVCCTCYCRGRNRCYCRVCEPILGYYPYCYOGGRVIC 300
DB 241 EELPINDYNTENGIEFDPMLDERGVCCTCYCRGRNRCYCRVCEPILGYYPYCYOGGRVIC 300
QY 301 RVIMPCKMVMYARMLGRV 317
DB 301 RVIMPCKMVMYARMLGRV 317

```

```

RESULT 11
US-10-063-616-116
Sequence 116, Application US/10063616
Publication No. US20030013855A1
GENERAL INFORMATION:
APPLICANT: Baton, Dan L.
APPLICANT: Filvaroff, Ellen
APPLICANT: Gerviloff, Mary E.
APPLICANT: Goddard, Audrey
APPLICANT: Goddard, Paul J.
APPLICANT: Grimaldi, Christopher J.
APPLICANT: Gurney, Austin L.
APPLICANT: Watanabe, Colin K.
TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
FILE REFERENCE: P3230R1C1
CURRENT APPLICATION NUMBER: US/10/063,616
CURRENT FILING DATE: 2002-05-03
Prior Application removed - See File Wrapper or Palm
NUMBER OF SEQ ID NOS: 170
SEQ ID NO 116
LENGTH: 317
TYPE: PRT
ORGANISM: Homo Sapien
US-10-063-616-116

```

```

Query Match 100.0%; Score 1745; DB 14; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
1 MAKNPENCDCIILNAEAFKSKIKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
1 MAKNPENCDCIILNAEAFKSKIKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
1 MAKNPENCDCIILNAEAFKSKIKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
QY 61 AYDMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLBVDHFKNGYTGIVFGLQKC 120
DB 61 AYDMEHTFYSNGEKKIYMEIDPVTRTEIFRSNGTDETLBVDHFKNGYTGIVFGLQKC 120
QY 121 FIKTOIVIPFSPSEEBEIDENEITTTFFEOQYIWPAPKPIENRFLKNSKILEICDN 180
DB 121 FIKTOIVIPFSPSEEBEIDENEITTTFFEOQYIWPAPKPIENRFLKNSKILEICDN 180
QY 181 VTWYINPTLISVSELODFEPEGEDLHFPANEKKGIEQNEQWVVPQYKVEKTRHARQASE 240
DB 181 VTWYINPTLISVSELODFEPEGEDLHFPANEKKGIEQNEQWVVPQYKVEKTRHARQASE 240
QY 241 EELPINDYNTENGIEFDPMLDERGVCCTCYCRGRNRCYCRVCEPILGYYPYCYOGGRVIC 300
DB 241 EELPINDYNTENGIEFDPMLDERGVCCTCYCRGRNRCYCRVCEPILGYYPYCYOGGRVIC 300

```

QY 301 RVIMPCNMWVARMIGRV 317  
DB 301 RVIMPCNMWVARMIGRV 317

## RESULT 12

US-10-121-049-524  
Sequence 524, Application US/10121049  
Publication No. US2003002239A1

## GENERAL INFORMATION:

APPLICANT: Baker, Kevin P.  
APPLICANT: Beresini, Maureen  
APPLICANT: Deforge, Laura  
APPLICANT: Desnoyers, Luc  
APPLICANT: Filvaroff, Ellen  
APPLICANT: Gao, Wei-Qiang  
APPLICANT: Gerritsen, Mary E.  
APPLICANT: Goddard, Audrey  
APPLICANT: Godowski, Paul J.  
APPLICANT: Gurney, Austin L.  
APPLICANT: Sherwood, Steven  
APPLICANT: Smith, Victoria  
APPLICANT: Stewart, Timothy A.  
APPLICANT: Tumas, Daniel  
APPLICANT: Watanabe, Colin K  
APPLICANT: Wood, William  
APPLICANT: Zhang, Zemin  
TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC  
FILE REFERENCE: P3330R1C17  
CURRENT APPLICATION NUMBER: US/10/121,049  
Prior Application removed - See File Wrapper or Palm  
NUMBER OF SEQ ID NOS: 550  
LENGTH: 317  
TYPE: PRT  
ORGANISM: Homo Sapien  
US-10-121-049-524

Query Match 100.0%; Score 1745; DB 14; Length 317;  
Best Local Similarity 100.0%; Pred. No. 2.5e-156;

Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALTLIVLFMSKHFMPVPPKX 60  
DB 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALTLIVLFMSKHFMPVPPKX 60  
QY 61 AYMEHTFVSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
DB 61 AYMEHTFVSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
QY 121 FIKTQIKVIFPESPEEIDENEITTTTFEQSVIWPAPKPIENRDFLNSKILEICDN 180  
DB 121 FIKTQIKVIFPESPEEIDENEITTTTFEQSVIWPAPKPIENRDFLNSKILEICDN 180  
QY 181 VTMWYNPTLISVSELQDFEEEGEDLHFPANEKKGIEQNEQWVVPQVKEKTRHARQASE 240  
DB 181 VTMWYNPTLISVSELQDFEEEGEDLHFPANEKKGIEQNEQWVVPQVKEKTRHARQASE 240  
QY 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRYCRVCEPLLGYYPPYCYOGGRVYC 300  
DB 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRYCRVCEPLLGYYPPYCYOGGRVYC 300  
QY 301 RVIMPCNMWVARMIGRV 317  
DB 301 RVIMPCNMWVARMIGRV 317

## RESULT 13

US-10-123-904-524  
Sequence 524, Application US/10123904  
Publication No. US20030022328A1

GENERAL INFORMATION:  
APPLICANT: Baker, Kevin P.  
APPLICANT: Beresini, Maureen  
APPLICANT: Deforge, Laura  
APPLICANT: Desnoyers, Luc  
APPLICANT: Filvaroff, Ellen  
APPLICANT: Gao, Wei-Qiang  
APPLICANT: Gerritsen, Mary E.  
APPLICANT: Goddard, Audrey  
APPLICANT: Godowski, Paul J.  
APPLICANT: Gurney, Austin L.  
APPLICANT: Sherwood, Steven  
APPLICANT: Smith, Victoria  
APPLICANT: Stewart, Timothy A.  
APPLICANT: Tumas, Daniel  
APPLICANT: Watanabe, Colin K  
APPLICANT: Wood, William  
APPLICANT: Zhang, Zemin

TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC  
FILE REFERENCE: P3330R1C54  
CURRENT APPLICATION NUMBER: US/10/123,904  
Prior Application removed - See File Wrapper or Palm  
NUMBER OF SEQ ID NOS: 550  
SEQ ID NO 524  
LENGTH: 317  
TYPE: PRT  
ORGANISM: Homo Sapien  
US-10-123-904-524

## Query Match

100.0%; Score 1745; DB 14; Length 317;  
Best Local Similarity 100.0%; Pred. No. 2.5e-156;  
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALTLIVLFMSKHFMPVPPKX 60  
DB 1 MAKNPENCDCCHILNAEAFSKKIKCSLKICGLVFGIALTLIVLFMSKHFMPVPPKX 60  
QY 61 AYMEHTFVSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
DB 61 AYMEHTFVSNGEKKKIYMEIDPVTRTEIFRSNGTDETLVHDFKNGYTGIVFGLQKC 120  
QY 121 FIKTQIKVIFPESPEEIDENEITTTTFEQSVIWPAPKPIENRDFLNSKILEICDN 180  
DB 121 FIKTQIKVIFPESPEEIDENEITTTTFEQSVIWPAPKPIENRDFLNSKILEICDN 180  
QY 181 VTMWYNPTLISVSELQDFEEEGEDLHFPANEKKGIEQNEQWVVPQVKEKTRHARQASE 240  
DB 181 VTMWYNPTLISVSELQDFEEEGEDLHFPANEKKGIEQNEQWVVPQVKEKTRHARQASE 240  
QY 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRYCRVCEPLLGYYPPYCYOGGRVYC 300  
DB 241 EELPINDYENGIEFPDMLDERGYCCICRRGNRYCRVCEPLLGYYPPYCYOGGRVYC 300  
QY 301 RVIMPCNMWVARMIGRV 317  
DB 301 RVIMPCNMWVARMIGRV 317

## RESULT 14

US-10-140-470-524  
Sequence 524, Application US/10140470  
Publication No. US20030022331A1

## GENERAL INFORMATION:

APPLICANT: Baker, Kevin P.  
APPLICANT: Beresini, Maureen  
APPLICANT: Deforge, Laura  
APPLICANT: Desnoyers, Luc  
APPLICANT: Filvaroff, Ellen  
APPLICANT: Gao, Wei-Qiang  
APPLICANT: Gerritsen, Mary E.  
APPLICANT: Goddard, Audrey

```

; APPLICANT: Godowski, Paul J.
; APPLICANT: Gurney, Austin L.
; APPLICANT: Sherwood, Steven
; APPLICANT: Smith, Victoria
; APPLICANT: Stewart, Timothy A.
; APPLICANT: Tamas, Daniel
; APPLICANT: Watanabe, Colin K
; APPLICANT: Wood, William
; APPLICANT: Zhang, Zemin
; TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
; FILE REFERENCE: P330R1C160
; CURRENT APPLICATION NUMBER: US/10/140,470
; PRIORITY FILING DATE: 2002-05-06
; PRIOR APPLICATION removed - See Palm or File Wrapper
; NUMBER OF SEQ ID NOS: 550
; SEQ ID NO 524
; LENGTH: 317
; TYPE: PRT
; ORGANISM: Homo Sapien
US-10-140-470-524

```

```

Query Match      100.0%; Score 1745; DB 14; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

```

```

QY 1 MAKNPENCDCHILNAEAFKSKKICKSLKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
DB 1 MAKNPENCDCHILNAEAFKSKKICKSLKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSGNGTDETLVHDFKNGYTGIFVGLQKC 120
DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSGNGTDETLVHDFKNGYTGIFVGLQKC 120
QY 121 FIKTQIKVIFPESPEEEDIDENEITTFPQSVIWPAPKPIENRDLKNSKILEICDN 180
DB 121 FIKTQIKVIFPESPEEEDIDENEITTFPQSVIWPAPKPIENRDLKNSKILEICDN 180
QY 181 VTMWINPTLISVSELDQFEEEGEDLHFPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
DB 181 VTMWINPTLISVSELDQFEEEGEDLHFPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
QY 241 EELPINDYENGIERPDMIDERGYCCICRGRNRCRVCCEPLIGYPPYCYOGGRVIC 300
DB 241 EELPINDYENGIERPDMIDERGYCCICRGRNRCRVCCEPLIGYPPYCYOGGRVIC 300
QY 301 RVIMPCNMWVARMIGRV 317
DB 301 RVIMPCNMWVARMIGRV 317

```

```

RESULT 15
US-10-063-502-116
; Sequence 116, Application US/10063502
; Publication No. US20030023042A1
; GENERAL INFORMATION:
; APPLICANT: Eaton, Dan L.
; APPLICANT: Filvaroff, Ellen
; APPLICANT: Gettlesen, Mary E.
; APPLICANT: Goddard, Audrey
; APPLICANT: Godowski, Paul J.
; APPLICANT: Grimaldi, Christopher J.
; APPLICANT: Gurney, Austin L.
; APPLICANT: Watanabe, Colin K.
; APPLICANT: Wood, William I.
; TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
; FILE REFERENCE: P3230R1C1
; CURRENT APPLICATION NUMBER: US/10/063,502
; PRIORITY FILING DATE: 2002-05-01
; PRIOR APPLICATION removed - See File Wrapper or Palm
; NUMBER OF SEQ ID NOS: 170
; SEQ ID NO 116

```

```

; LENGTH: 317
; TYPE: PRT
; ORGANISM: Homo Sapien
US-10-063-502-116

```

```

Query Match      100.0%; Score 1745; DB 14; Length 317;
Best Local Similarity 100.0%; Pred. No. 2.5e-156;
Matches 317; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

```

```

QY 1 MAKNPENCDCHILNAEAFKSKKICKSLKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
DB 1 MAKNPENCDCHILNAEAFKSKKICKSLKICGLVFGILATLTLVLFWSGKHFWPEVPRK 60
QY 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSGNGTDETLVHDFKNGYTGIFVGLQKC 120
DB 61 AYDMEHTFYSNGEKKKIYMEIDPVTRTEIFRSGNGTDETLVHDFKNGYTGIFVGLQKC 120
QY 121 FIKTQIKVIFPESPEEEDIDENEITTFPQSVIWPAPKPIENRDLKNSKILEICDN 180
DB 121 FIKTQIKVIFPESPEEEDIDENEITTFPQSVIWPAPKPIENRDLKNSKILEICDN 180
QY 181 VTMWINPTLISVSELDQFEEEGEDLHFPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
DB 181 VTMWINPTLISVSELDQFEEEGEDLHFPANEKKGIEQNEQWVPQVKEKTRHARQASE 240
QY 241 EELPINDYENGIERPDMIDERGYCCICRGRNRCRVCCEPLIGYPPYCYOGGRVIC 300
DB 241 EELPINDYENGIERPDMIDERGYCCICRGRNRCRVCCEPLIGYPPYCYOGGRVIC 300
QY 301 RVIMPCNMWVARMIGRV 317
DB 301 RVIMPCNMWVARMIGRV 317

```

```

Search completed: March 30, 2004, 14:38:19
Job time : 46 secs

```